# (19) World Intellectual Property Organization International Bureau





# (43) International Publication Date 9 August 2001 (09.08.2001)

# **PCT**

# (10) International Publication Number WO 01/57194 A2

(51) International Patent Classification<sup>7</sup>: C12N 9/00

(21) International Application Number: PCT/US01/03471

(22) International Filing Date: 2 February 2001 (02.02.2001)

(25) Filing Language: English

(26) Publication Language: English

(30) Priority Data:

60/179,982	3 February 2000 (03.02.2000)	US
60/183,542	18 February 2000 (18.02.2000)	US
60/213,124	22 June 2000 (22.06.2000)	US
60/220,970	26 July 2000 (26.07.2000)	US
09/657,986	8 September 2000 (08.09.2000)	US
60/234,840	22 September 2000 (22.09.2000)	US

(63) Related by continuation (CON) or continuation-in-part (CIP) to earlier applications:

60/179,982 (CIP)
3 February 2000 (03.02.2000)
60/183,542 (CIP)
18 February 2000 (18.02.2000)
60/213,124 (CIP)
22 June 2000 (22.06.2000)
60/220,970 (CIP)
26 July 2000 (26.07.2000)
09/657,986 (CIP)
8 September 2000 (08.09.2000)
60/234,840 (CIP)
22 September 2000 (22.09.2000)

(71) Applicant (for all designated States except US): CORVAS INTERNATIONAL, INC. [US/US]; 3030 Science Park Road, San Diego, CA 92121 (US).

- (72) Inventors; and
- (75) Inventors/Applicants (for US only): MADISON, Edwin, L. [US/US]; 501 Stratford Court, Del Mar, CA 92014 (US). ONG, Edgar, O. [CA/US]; 8328 Regents Road #3C, San Diego, CA 92122 (US). YEH, Jiunn-chern [—/US]; 3128-E Via Alicante Drive, La Jolla, CA 92037 (US).
- (74) Agents: SEIDMAN, Stephanie, L. et al.; Heller Ehrman White & McAuliffe LLP, Suite 700, 4250 Executive Square, La Jolla, CA 92037 (US).
- (81) Designated States (national): AE, AG, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

# Published:

 without international search report and to be republished upon receipt of that report

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.



1/57194 A

**(54) Title:** NUCLEIC ACID MOLECULES ENCODING TRANSMEMBRANE SERINE PROTEASES, THE ENCODED PROTEINS AND METHODS BASED THEREON

-1-

# NUCLEIC ACID MOLECULES ENCODING TRANSMEMBRANE SERINE PROTEASES, THE ENCODED PROTEINS AND METHODS BASED THEREON RELATED APPLICATIONS

Benefit of priority is claimed to U.S. provisional application Serial No. 60/179,982, to Edwin L. Madison and Edgar O. Ong, filed February 3, 2000, 5 entitled "NUCLEOTIDE AND PROTEIN SEQUENCES OF A TRANSMEMBRANE SERINE PROTEASE AND METHODS BASED THEREOF"; to U.S. provisional application Serial No. 60/183,542, to Edwin L. Madison and Edgar O. Ong, filed February 18, 2000, entitled "NUCLEOTIDE AND PROTEIN SEQUENCES OF A TRANSMEMBRANE SERINE PROTEASE AND METHODS BASED THEREOF"; to 10 U.S. provisional application Serial No. 60/213,124, to Edwin L. Madison and Edgar O. Ong, filed June 22, 2000, entitled "NUCLEOTIDE AND PROTEIN SEQUENCES OF A TRANSMEMBRANE SERINE PROTEASE AND METHODS BASED THEREOF"; to U.S. provisional application Serial No. 60/220,970, to Edwin L. Madison and Edgar O. Ong, filed July 26, 2000, entitled 15 "NUCLEOTIDE AND PROTEIN SEQUENCES OF A TRANSMEMBRANE SERINE PROTEASE AND METHODS BASED THEREOF"; and to U.S. provisional application Serial No. 60/234,840 to Edwin L. Madison, Edgar O. Ong and Jiunn-Chern Yeh, filed September 22, 2000, entitled "NUCLEIC ACID MOLECULES 20 ENCODING TRANSMEMBRANE SERINE PROTEASES, THE ENCODED PROTEINS AND METHODS BASED THEREON" is claimed herein. Benefit of priority is also claimed to U.S. application Serial No. 09/657,968, to Edwin L. Madison, Joseph Edward Semple, Gary Samuel Coombs, John Eugene Reiner, Edgar O. Ong, Gian Luca Araldi, filed September 8, 2000, entitled "INHIBITORS OF SERINE PROTEASE ACTIVITY OF MATRIPTASE OR MTSP1". This application is a 25 continuation-in-part of U.S. application Serial No. 09/657,986.

This application is related to U.S. provisional application Serial No. 60/166,391 to Edwin L. Madison and Edgar O. Ong, filed November 18, 1999 entitled "NUCLEOTIDE AND PROTEIN SEQUENCES OF PROTEASE DOMAINS OF ENDOTHELIASE AND METHODS BASED THEREON". This a application is also

30

-2-

related to International PCT application No. PCT/US00/31803, filed November 17, 2000.

Where permitted, the above-noted provisional applications, patent application and International PCT application are incorporated by reference in their entirety. All patents, applications, published applications and other publications and sequences from GenBank and other data bases referred to herein are incorporated by reference in their entirety.

# FIELD OF INVENTION

10

20

25

30

Nucleic acid molecules that encode proteases and portions thereof, particularly protease domains are provided. Also provided are prognostic, diagnostic and therapeutic methods using the proteases and domains thereof and the encoding nucleic acid molecules.

#### **BACKGROUND OF THE INVENTION AND OBJECTS THEREOF**

Cancer a leading cause of death in the United States, developing in one in three Americans; one of every four Americans dies of cancer. Cancer is characterized by an increase in the number of abnormal neoplastic cells, which proliferate to form a tumor mass, the invasion of adjacent tissues by these neoplastic tumor cells, and the generation of malignant cells that metastasize via the blood or lymphatic system to regional lymph nodes and to distant sites.

Among the hallmarks of cancer is a breakdown in the communication among tumor cells and their environment. Normal cells do not divide in the absence of stimulatory signals, cease dividing in the presence of inhibitory signals. Growth-stimulatory and growth-inhibitory signals, are routinely exchanged between cells within a tissue. In a cancerous, or neoplastic, state, a cell acquires the ability to "override" these signals and to proliferate under conditions in which normal cells do not grow.

In order to proliferate tumor cells acquire a number of distinct aberrant traits reflecting genetic alterations. The genomes of certain well-studied tumors carry several different independently altered genes, including activated oncogenes and inactivated tumor suppressor genes. Each of these genetic changes appears to be responsible for imparting some of the traits that, in the aggregate, represent the full neoplastic phenotype.

-3-

A variety of biochemical factors have been associated with different phases of metastasis. Cell surface receptors for collagen, glycoproteins such as laminin, and proteoglycans, facilitate tumor cell attachment, an important step in invasion and metastases. Attachment triggers the release of degradative enzymes which facilitate the penetration of tumor cells through tissue barriers. Once the tumor cells have entered the target tissue, specific growth factors are required for further proliferation. Tumor invasion (or progression) involves a complex series of events, in which tumor cells detach from the primary tumor, break down the normal tissue surrounding it, and migrate into a blood or lymphatic vessel to be carried to a distant site. The breaking down of normal tissue barriers is accomplished by the elaboration of specific enzymes that degrade the proteins of the extracellular matrix that make up basement membranes and stromal components of tissues.

10

15

20

25

30

A class of extracellular matrix degrading enzymes have been implicated in tumor invasion. Among these are the matrix metalloproteinases (MMP). For example, the production of the matrix metalloproteinase stromelysin is associated with malignant tumors with metastatic potential (see, e.g., McDonnell et al. (1990) Smnrs. in Cancer Biology 1:107-115; McDonnell et al. (1990) Cancer and Metastasis Reviews 9:309-319).

The capacity of cancer cells to metastasize and invade tissue is facilitated by degradation of the basement membrane. Several proteinase enzymes, including the MMPs, have been reported to facilitate the process of invasion of tumor cells. MMPs are reported to enhance degradation of the basement membrane, which thereby permits tumorous cells to invade tissues. For example, two major metalloproteinases having molecular weights of about 70 kDa and 92 kDa appear to enhance ability of tumor cells to metastasize.

# Type II Transmembrane Serine Proteases (TTSPs)

In addition to the MMPs, serine proteases have been implicated in neoplastic disease progression. Most serine proteases, which are either secreted enzymes or are sequestered in cytoplasmic storage organelles, have roles in blood coagulation, wound healing, digestion, immune responses and tumor invasion and metastasis. A class cell surface proteins designated type II

WO 01/57194

30

PCT/US01/03471

transmembrane serine proteases, which are membrane-anchored proteins with N-terminal extracellular domains, has been identified. As cell surface proteins, they are positioned to play a role in intracellular signal transduction and in mediating cell surface proteolytic events.

5 Cell surface proteolysis is a mechanism for the generation of biologically active proteins that mediate a variety of cellular functions. These membraneanchored proteins, include a disintegrin-like and metalloproteinase (ADAM) and membrane-type matrix metalloproteinase (MT-MMP). In mammals, at least 17 members of the family are known, including seven in humans (see, Hooper et al. (2001) J. Biol. Chem. 276:857-860). These include: corin (accession nos. 10 AF133845 and AB013874; see, Yan et al. (1999) J. Biol. Chem. 274:14926-14938; Tomia et al. (1998) J. Biochem. 124:784-789; Uan et al. (2000) Proc. Natl. Acad. Sci. U.S.A. 97:8525-8529); enterpeptidase (also designated enterokinase; accession no. U09860 for the human protein; see, Kitamoto et al. (1995) Biochem. 27: 4562-4568; Yahagi et al. (1996) Biochem. Biophys. Res. 15 Commun. 219:806-812; Kitamoto et al. (1994) Proc. Natl. Acad. Sci. U.S.A. 91:7588-7592; Matsushima et al. (1994) J. Biol. Chem. 269:19976-19982;); human airway trypsin-like protease (HAT; accession no. AB002134; see Yamaoka et al. J. Biol. Chem. 273:11894-11901); MTSP1 and matriptase (also 20 called TADG-15; see SEQ ID Nos. 1 and 2; accession nos. AF133086/AF118224, AF04280022; Takeuchi et al. (1999) Proc. Natl. Acad. Sci. U.S.A. 96:11054-1161; Lin et al. (1999) J. Biol. Chem. 274:18231-18236; Takeuchi et al. (2000) J. Biol. Chem. 275:26333-26342; and Kim et al. (1999) Immunogenetics 49:420-429); hepsin (see, accession nos. M18930, AF030065, X70900; Leytus et al. (1988) Biochem. 27: 11895-11901; Vu et al. (1997) J. 25 Biol. Chem. 272:31315-31320; and Farley et al. (1993) Biochem. Biophys. Acta 1173:350-352; and see, U.S. Patent No. 5,972,616); TMPRS2 (see, Accession Nos. U75329 and AF113596; Paoloni-Giacobino et al. (1997) Genomics 44:309-320; and Jacquinet et al. (2000) FEBS Lett. 468: 93-100); and TMPRSS4 (see,

Serine proteases, including transmembrane serine proteases, have been implicated in processes involved in neoplastic development and progression.

Accession No. NM 016425; Wallrapp et al. (2000) Cancer 60:2602-2606).

-5-

While the precise role of these proteases has not been elaborated, serine proteases and inhibitors thereof are involved in the control of many intra- and extracellular physiological processes, including degradative actions in cancer cell invasion, metastatic spread, and neovascularization of tumors, that are involved in tumor progression. It is believed that proteases are involved in the degradation of extracellular matrix (ECM) and contribute to tissue remodeling, and are necessary for cancer invasion and metastasis. The activity and/or expression of some proteases have been shown to correlate with tumor progression and development.

For example, a membrane-type serine protease MTSP1 (also called matriptase; see SEQ ID Nos. 1 and 2 from U.S. Patent No. 5,972,616; and GenBank Accession No. AF118224; (1999) *J. Biol. Chem. 274*:18231-18236; U.S. Patent No. 5,792,616; see, also Takeuchi (1999) *Proc. Natl. Acad. Sci. U.S.A. 96*:11054-1161) that is expressed in epithelial cancer and normal tissue (Takeucuhi *et al.* (1999) *Proc. Natl. Acad. Sci. USA*, 96(20):11054-61) has been identified. Matriptase was originally identified in human breast cancer cells as a major gelatinase (see, U.S. Patent No. 5,482,848), a type of matrix metalloprotease (MMP). It has been proposed that it plays a role in the metastasis of breast cancer. Its primary cleavage specificity is Arg-Lys residues.
Matriptase also is expressed in a variety of epithelial tissues with high levels of activity and/or expression in the human gastrointestinal tract and the prostate.

Prostate-specific antigen (PSA), a kallikrein-like serine protease, degrades extracellular matrix glycoproteins fibronectin and laminin, and, has been postulated to facilitate invasion by prostate cancer cells (Webber *et al.* (1995) *Clin. Cancer Res.*, 1(10):1089-94). Blocking PSA proteolytic activity with PSA-specific monoclonal antibodies results in a dose-dependent decrease *in vitro* in the invasion of the reconstituted basement membrane Matrigel by LNCaP human prostate carcinoma cells which secrete high levels of PSA.

25

30

Hepsin, a cell surface serine protease identified in hepatoma cells, is overexpressed in ovarian cancer (Tanimoto *et al.* (1997) *Cancer Res.*, 57(14):2884-7). The hepsin transcript appears to be abundant in carcinoma tissue and is almost never expressed in normal adult tissue, including normal

-6-

ovary. It has been suggested that hepsin is frequently overexpressed in ovarian tumors and therefore may be a candidate protease in the invasive process and growth capacity of ovarian tumor cells.

A serine protease-like gene, designated normal epithelial cell-specific 1 (NES1) (Liu et al., *Cancer Res.*, <u>56(14)</u>:3371-9 (1996)) has been identified. Although expression of the NES1 mRNA is observed in all normal and immortalized nontumorigenic epithelial cell lines, the majority of human breast cancer cell lines show a drastic reduction or a complete lack of its expression. The structural similarity of NES1 to polypeptides known to regulate growth factor activity and a negative correlation of NES1 expression with breast oncogenesis suggest a direct or indirect role for this protease-like gene product in the suppression of tumorigenesis.

Hence transmembrane serine proteases appear to be involved in the etiology and pathogenesis of tumors. There is a need to further elucidate their role in these processes and to identify additional transmembrane proteases. Therefore, it is an object herein to provide transmembrane serine protease (MTSP) proteins and nucleic acids encoding such MTSP proteases that are involved in the regulation of or participate in tumorigenesis and/or carcinogenesis. It is also an object herein to provide prognostic, diagnostic, therapeutic screening methods using the such proteases and the nucleic acids encoding such proteases.

# **SUMMARY OF THE INVENTION**

10

15

20

25

30

Provided herein are isolated protease domains of the Transmembrane Serine Protease family, particularly the Type II Transmembrane Serine Protease (TTSP) family (also referred to herein as MTSPs), and more particularly TTSP family members whose functional activity differs in tumor cells from non-tumor cells in the same tissue. For example, the MTSPs include those that are activated and/or expressed in tumor cells at different levels, typically higher, from non-tumor cells; and those from cells in which substrates therefor differ in tumor cells from non-tumor cells or otherwise alter the specificity of the MTSP.

The MTSP family as intended herein does not include any membrane anchored or spanning proteases that are expressed on endothelial cells.

-7-

Included among the MTSPs are several heretofore unidentified MTSP family members, designated herein as MTSP3 and MTSP4 and a new form of a protein designated herein as MTSP6. In addition to the protease domains of each of MTSP3 and MTSP4, the full-length proteins, including those that results from splice variants, zymogens and activated forms, and uses thereof, are also provided.

The protease domains as provided herein are single-chain polypeptides, with an N-terminus (such as IV, VV, IL and II) generated at the cleavage site (generally having the consensus sequence R\ddot VVGG, R\ddot IVGG, R\ddot IVNG, R\ddot ILGG, R\ddot VGLL, R\ddot ILGG or a variation thereof; an N-terminus R\ddot V or R\ddot I, where the arrow represents the cleavage point) when the zymogen is activated. To identify the protease domain an RI should be identified, and then the following amino acids compared to the above noted motif.

The protease domains generated herein, however, do not result from

10

activation, which produces a two chain activated product, but rather are single 15 chain polypeptides with the N-terminus include the consensus sequence ↓VVGG, ↓IVGG, ↓VGLL, ↓ILGG or ↓IVNG or other such motif at the Nterminus. As shown herein, such polypeptides, although not the result of activation and not double-chain forms, exhibit proteolytic (catalytic) activity. 20 These protease domain polypeptides are used in assays to screen for agents that modulate the activity of the MTSP. Such assays are also provided herein. In exemplary assays, the affects of test compounds in the ability of a protease domains to proteolytically cleave a known substrate, typically a fluorescently, chromogenically or otherwise detectably labeled substrate, are assessed. 25 Agents, generally compounds, particularly small molecules, that modulate the activity of the protease domain are candidate compounds for modulating the activity of the MTSP. The protease domains can also be used to produce singlechain protease-specific antibodies. The protease domains provided herein include, but are not limited to, the single chain region having an N-terminus at 30 the cleavage site for activation of the zymogen, through the C-terminus, or Cterminal truncated portions thereof that exhibit proteolytic activity as a singlechain polypeptide in in vitro proteolysis assays, of any MTSP family member,

-8-

preferably from a mammal, including and most preferably human, that, for example, is expressed in tumor cells at different levels from non-tumor cells, and that is not expressed on an endothelial cell. These include, but are not limited to : MTSP1 (or matriptase), MTSP3, MTSP4 and MTSP6. Other MTSP protease 5 domains of interest herein, particularly for use in in vitro drug screening proteolytic assays, include, but are not limited to: corin (accession nos. AF133845 and AB013874; see, Yan et al. (1999) J. Biol. Chem. 274:14926-14938; Tomia et al. (1998) J. Biochem. 124:784-789; Uan et al. (2000) Proc. Natl. Acad. Sci. U.S.A. 97:8525-8529; SEQ ID Nos. 61 and 62 for the human 10 protein); enterpeptidase (also designated enterokinase; accession no. U09860 for the human protein; see, Kitamoto et al. (1995) Biochem. 27: 4562-4568; Yahagi et al. (1996) Biochem. Biophys. Res. Commun. 219:806-812; Kitamoto et al. (1994) Proc. Natl. Acad. Sci. U.S.A. 91:7588-7592; Matsushima et al. (1994) J. Biol. Chem. 269:19976-19982; see SEQ ID Nos. 63 and 64 for the human protein); human airway trypsin-like protease (HAT; accession no. 15 AB002134; see Yamaoka et al. J. Biol. Chem. 273:11894-11901; SEQ ID Nos. 65 and 66 for the human protein); hepsin (see, accession nos. M18930, AF030065, X70900; Leytus et al. (1988) Biochem. 27: 11895-11901; Vu et al. (1997) J. Biol. Chem. 272:31315-31320; and Farley et al. (1993) Biochem. 20 Biophys. Acta 1173:350-352; SEQ ID Nos. 67 and 68 for the human protein); TMPRS2 (see, Accession Nos. U75329 and AF113596; Paoloni-Giacobino et al. (1997) Genomics 44:309-320; and Jacquinet et al. (2000) FEBS Lett. 468: 93-100; SEQ ID Nos. 69 and 70 for the human protein) TMPRSS4 (see, Accession No. NM 016425; Wallrapp et al. (2000) Cancer 60:2602-2606; SEQ ID Nos. 71 25 and 72 for the human protein); and TADG-12 (also designated MTSP6, see SEQ ID Nos. 11 and 12; see International PCT application No. WO 00/52044, which

Also provided are muteins of the single chain protease domains and MTSPs, particularly muteins in which the Cys residue in the protease domain that is free (*i.e.*, does not form disulfide linkages with any other Cys residue in the protein) is substituted with another amino acid substitution, preferably with a conservative amino acid substitution or a substitution that does not eliminate the

claims priority to U.S. application Serial No. 09/261,416).

30

-9-

activity, and muteins in which a glycosylation site(s) is eliminated. Muteins in which other conservative amino acid substitutions in which catalytic activity is retained are also contemplated (see, *e.g.*, Table 1, for exemplary amino acid substitutions). See, also, Figure 4, which identifies the free Cys residues in MTSP3, MTSP4 and MTSP6.

5

10

15

20

25

30

Hence, provided herein are members of a family of transmembrane serine protease (MTSP) proteins, and functional domains, especially protease (or catalytic) domains thereof, muteins and other derivatives and analogs thereof. Also provided herein are nucleic acids encoding the MTSPs.

Exemplary MTSPs (see, e.g., SEQ ID No. 1-12, 49 and 50) are provided herein, as are the single chain protease domains thereof as follows: SEQ ID Nos. 1, 2, 49 and 50 set forth amino acid and nucleic acid sequences of MTSP1 and the protease domain thereof; SEQ ID No. 3 sets forth the MTSP3 nucleic acid sequence and SEQ ID No. 4 the encoded MTSP3 amino acids; SEQ ID No. 5 MTSP4 a nucleic acid sequence of the protease domain and SEQ ID No. 6 the encoded MTSP4 amino acid protease domain; SEQ ID No. 7 MTSP4-L a nucleic acid sequence and SEQ ID No. 8 the encoded MTSP4-L amino acid sequence; SEQ ID No. 9 an MTSP4-S encoding nucleic acid sequence and SEQ ID No. 10 the encoded MTSP4-S amino acid sequence; and SEQ ID No. 11 an MTSP6 encoding nucleic acid sequence. The single chain protease domains of each are delineated below.

Nucleic acid molecules that encode a single-chain protease domain or catalytically active portion thereof are provided. Also provided are nucleic acid molecules that hybridize to such *MTSP* encoding nucleic acid along their full length and encode the protease domain or portion thereof are provided. Hybridization is preferably effected under conditions of at least low, generally at least moderate, and often high stringency.

Additionally provided herein are antibodies that specifically bind to the MTSPs, cells, combinations, kits and articles of manufacture that contain the nucleic acid encoding the MTSP and/or the MTSP. Further provided herein are prognostic, diagnostic, therapeutic screening methods using MTSPs and the nucleic acids encoding MTSP. Also provided are transgenic non-human animals

-10-

bearing inactivated genes encoding the MTSP and bearing the genes encoding the MTSP under non-native promotor control. Such animals are useful in animal models of tumor initation, growth and/or progression models.

Provided herein are members of a family of membrane serine proteases (MTSP) that are expressed in certain tumor or cancer cells such lung, prostate, 5 colon and breast cancers. In particular, it is shown herein, that MTSPs, particularly, MTSP3, MTSP4 and MTSP6 are expressed in lung carcinoma, breast carcinoma, colon adenocarcinoma and/or ovarian carcinomas as well as in certain normal cells and tissues (see e.g., EXAMPLES for tissue-specific expression profiles of each protein exemplified herein). The MTSPs that are of 10 particular interest herein, are those that are expressed in tumor cells, for example, those that appear to be expressed at different levels in tumor cells from normal cells, or whose functional activity is different in tumor cells from normal cells, such as by an alteration in a substrate therefor, or a cofactor. Hence the MTSP provided herein can serve as diagnostic markers for certain 15 tumors. The level of activated MTSP3, MTSP4 and MTSP6 can be diagnostic of prostate cancer. In addition, MTSP4 is expressed and/or activated in lymphomas, leukemias, lung cancer, breast, prostrate and colon cancers. MTSP6 is activated and/or expressed in breast, lung, prostate, colon and ovarian cancers. Furthermore, compounds that modulate the activity of these MTSPs, 20 as assessed by the assays provided herein, particularly the in vitro proteolytic assays that use the single chain protease domains, are potential therapeutic candidates for treatment of various malignancies and neoplastic disease.

Also provided herein are methods of modulating the activity of the MTSPs and screening for compounds that modulate, including inhibit, antagonize, agonize or otherwise alter the activity of the MTSPs. Of particular interest is the extracellular domain of these MTSPs that includes the proteolytic (catalytic) portion of the protein.

MTSP proteins, including, but not limited to, MTSP3, MTSP4, and MTSP6, including splice variants thereof, and nucleic acids encoding MTSPs, and domains, derivatives and analogs thereof are provided herein. Single chain protease domains, in the N-terminal is that which would be generated by

25

30

-11-

activation of the zymogen, from any MTSP, particularly those that are not expressed in endothelial cells and that are expressed in tumor cells are also provided.

Antibodies that specifically bind to the MTSP, particularly the single chain protease domain, and any and all forms of MTSP3 and MTSP4, and cells, combinations, kits and articles of manufacture containing the MTSP proteins, domains thereof, or encoding nucleic acids are also provided herein. Transgenic non-human animals bearing inactivated genes encoding the MTSP and bearing the genes encoding the MTSP under a non-native promotor control are additionally provided herein. Also provided are nucleic acid molecules encoding each of the MTSPs and domains thereof.

10

15

20

25

30

Also provided are plasmids containing any of the nucleic acid molecules provided herein. Cells containing the plasmids are also provided. Such cells include, but are not limited to, bacterial cells, yeast cells, fungal cells, plant cells, insect cells and animal cells.

Also provided is a method of producing a MTSP by growing the abovedescribed cells under conditions whereby the MTSP is expressed by the cells, and recovering the expressed MTSP protein. Methods for isolating nucleic acid encoding other MTSPs are also provided.

Also provided are cells, preferably eukaryotic cells, such as mammalian cells and yeast cells, in which the MTSP protein, preferably MTSP3 and MTSP4, is expressed in the surface of the cells. Such cells are used in drug screening assays to identify compounds that modulate the activity of the MTSP protein. These assays including *in vitro* binding assays, and transcription based assays in which signal transduction mediated by the MTSP is assessed.

Further provided herein are prognostic, diagnostic and therapeutic screening methods using the MTSP and the nucleic acids encoding MTSP. In particular, the prognostic, diagnostic and therapeutic screening methods are used for preventing, treating, or for finding agents useful in preventing or treating, tumors or cancers such as lung carcinoma, colon adenocarcinoma and ovarian carcinoma.

Also provided are methods for screening for compounds that modulate the activity of any MTSP. The compounds are identified by contacting them with the MTSP and a substrate for the MTSP. A change in the amount of substrate cleaved in the presence of the compounds compared to that in the absence of the compound indicates that the compound modulates the activity of the MTSP. Such compounds are selected for further analyses or for use to modulate the activity of the MTSP, such as inhibitors or agonists. The compounds can also be identified by contacting the substrates with a cell that expresses the MTSP or the extracellular domain or proteolytically active portion thereof. For assays in which the extracellular domain or a proteolytically active portion thereof is employed, the MTSP is any MTSP that is expressed on cells, other than endothelial cells, including, but not limited to MTSP1, MTSP3, MTSP4 and MTSP6.

10

15

20

25

30

Also provided herein are modulators of the activity of the MTSP, especially the modulators obtained according to the screening methods provide herein. Such modulators may have use in treating cancerous conditions, and other neoplastic conditions.

Pharmaceutical composition containing the protease domains of an MTSP protein, and the MTSP proteins, MTSP3, MTSP4 and MTSP6 are provided herein in a pharmaceutically acceptable carrier or excipient are provided herein.

Also provided are articles of manufacture that contain the MTSP proteins and protease domains of MTSPs in single chain form. The articles contain a) packaging material; b) the polypeptide (or encoding nucleic acid), particularly the single chain protease domain thereof; and c) a label indicating that the article is for using ins assays for identifying modulators of the activities of an MTSP protein is provided herein.

Conjugates containing a) a MTSP protease domain in single chain from; and b) a targeting agent linked to the MTSP directly or via a linker, wherein the agent facilitates: i) affinity isolation or purification of the conjugate; ii) attachment of the conjugate to a surface; iii) detection of the conjugate; or iv) targeted delivery to a selected tissue or cell, is provided herein. The conjugate

-13-

can contain a plurality of agents linked thereto. The conjugate can be a chemical conjugate; and it can be a fusion protein.

5

10

15

20

25

30

In yet another embodiment, the targeting agent is a protein or peptide fragment. The protein or peptide fragment can include a protein binding sequence, a nucleic acid binding sequence, a lipid binding sequence, a polysaccharide binding sequence, or a metal binding sequence.

Method of diagnosing a disease or disorder characterized by detecting an aberrant level of an MTSP, particularly an MTSP3, MTSP4 or MTSP 6, in a subject is provided. The method can be practiced by measuring the level of the DNA, RNA, protein or functional activity of the MTSP. An increase or decrease in the level of the DNA, RNA, protein or functional activity of the MTSP, relative to the level of the DNA, RNA, protein or functional activity found in an analogous sample not having the disease or disorder (or other suitable control) is indicative of the presence of the disease or disorder in the subject or other relative any other suitable control.

Combinations are provided herein. The combination can include: a) an inhibitor of the activity of an MTSP; and b) an anti-cancer treatment or agent. The MTSP inhibitor and the anti-cancer agent can be formulated in a single pharmaceutical composition or each is formulated in a separate pharmaceutical composition. The MTSP inhibitor can be an antibody or a fragment or binding portion thereof against the MTSP, such as an antibody that specifically binds to the protease domain, an inhibitor of the MTSP production, or an inhibitor of the MTSP membrane-localization or an inhibitor of MTSP activation. Other MTSP inhibitors include, but are not limited to, an antisense nucleic acid encoding the MTSP, particularly a portion of the protease domain; a nucleic acid encoding at least a portion of a gene encoding the MTSP with a heterologous nucleotide sequence inserted therein such that the heterologous sequence inactivates the biological activity encoded MTSP or the gene encoding it. The portion of the gene encoding the MTSP preferably flanks the heterologous sequence to promote homologous recombination with a genomic gene encoding the MTSP.

Also, provided are methods for treating or preventing a tumor or cancer in a mammal by administering to a mammal an effective amount of an inhibitor of

an MTSP3, MTSP4 or MTSP6, whereby the tumor or cancer is treated or prevented. The MTSP inhibitor used in the treatment or for prophylaxis is administered with a pharmaceutically acceptable carrier or excipient. The mammal treated can be a human. The treatment or prevention method can additionally include administering an anti-cancer treatment or agent simultaneously with or subsequently or before administration of the MTSP inhibitor.

Also provided is a recombinant non-human animal in which an endogenous gene of an MTSP has been deleted or inactivated by homologous recombination or insertional mutagenesis of the animal or an ancestor thereof. A recombinant non-human animal is provided herein, where the gene of an MTSP is under control of a promoter that is not the native promoter of the gene or that is not the native promoter of the gene in the non-human animal or where the nucleic acid encoding the MTSP is heterologous to the non-human animal and the promoter is the native or a non-native promoter.

Also provided are methods of treatments of tumors by administering a prodrug that is activated by an MTSP that is expressed or active in tumor cells, particularly those in which its functional activity in tumor cells is greater than in none-tumor cells. The prodrug is administered and, upon administration, active MTSP expressed on cells cleaves the prodrug and releases active drug in the vicinity of these cells. The active anti-cancer drug accumulates in the vicinity of the tumor. This is particularly useful in instances in which an MTSP is expressed or active in greater quantity, higher level or predominantly in tumor cells compared to other cells.

# 25 BRIEF DESCRIPTION OF DRAWINGS

10

15

20

30

Figure 1 illustrates the domain organization of the MTSP3;

Figure 2 illustrates the domain organization of the MTSP4 splice variants and domains thereof; MTSP4-L includes a transmembrane domain, a CUB domain, a low density lipoprotein receptor (LDLR) domains, and a serine protease catalytic domain; MTSP4-S lacking the portion between amino acids 136-279.

Figure 3 depicts the domain organization of MTSP6.

-15-

Figure 4 provides an alignment of the C-terminal portions of MTSP3, the two splice variant-encoded forms of MTSP4, and MTSP6, that encompasses the protease domains thereof; the figure shows the cleavage sites, which form the N-terminus of the protease domain of each protein; a potential glycosylation site is noted and the free Cys residues in the protease domain of each are noted (\*). Muteins of each protein may be prepared by replacing the residues in the glycosylation site, particularly the N residue, and the free Cys residues, with preferably conservative amino acid residues. Such muteins are also provided herein.

# 10 DETAILED DESCRIPTION OF THE INVENTION

#### A. DEFINITIONS

15

20

25

30

Unless defined otherwise, all technical and scientific terms used herein have the same meaning as is commonly understood by one of ordinary skill in the art to which this invention belongs. All patents, applications, published applications and other publications and sequences from GenBank and other data bases referred to herein are incorporated by reference in their entirety.

As used herein, the abbreviations for any protective groups, amino acids and other compounds, are, unless indicated otherwise, in accord with their common usage, recognized abbreviations, or the IUPAC-IUB Commission on Biochemical Nomenclature (see, (1972) <u>Biochem.</u> 11:942-944).

As used herein, serine protease refers to a diverse family of proteases wherein a serine residue is involved in the hydrolysis of proteins or peptides. The serine residue can be part of the catalytic triad mechanism, which includes a serine, a histidine and an aspartic acid in the catalysis, or be part of the hydroxyl/e-amine or hydroxyl/a-amine catalytic dyad mechanism, which involves a serine and a lysine in the catalysis.

As used herein, "transmembrane serine protease (MTSP)" refers to a family of transmembrane serine proteases that share common structural features as described herein (see, also Hooper *et al.* (2001) *J. Biol. Chem.276*:857-860). Thus, reference, for example, to "MTSP" encompasses all proteins encoded by the MTSP gene family, including but are not limited to: MTSP1, MTSP3, MTSP4 and MTSP6, or an equivalent molecule obtained from any other source or that

-16-

has been prepared synthetically or that exhibits the same activity. Other MTSPs include, but are not limited to, corin, enterpeptidase, human airway trypsin-like protease (HAT), MTSP1, TMPRSS2, and TMPRSS4. Sequences of encoding nucleic molecules and the encoded amino acid sequences of exemplary MTSPs and/or domains thereof are set forth in SEQ ID Nos. 1-12, 49, 50 and 61-72. The term also encompass MTSPs with conservative amino acid substitutions that do not substantially alter activity of each member, and also encompasses splice variants thereof. Suitable conservative substitutions of amino acids are known to those of skill in this art and may be made generally without altering the biological activity of the resulting molecule. Of particular interest are MTSPs of mammalian, including human, origin. Those of skill in this art recognize that, in general, single amino acid substitutions in non-essential regions of a polypeptide do not substantially alter biological activity (see, e.g., Watson et al. Molecular Biology of the Gene, 4th Edition, 1987, The Bejacmin/Cummings Pub. co., p.224).

10

15

20

25

30

As used herein, a "protease domain of an MTSP" refers to the protease domain of MTSP that is located within the extracellular domain of a MTSP and exhibits serine proteolytic activity. It includes at least the smallest fragment thereof that acts catalytically as a single chain form. Hence it is at least the minimal portion of the extracellular domain that exhibits proteolytic activity as assessed by standard assays *in vitro* assays. Those of skill in this art recognize that such protease domain is the portion of the protease that is structurally equivalent to the trypsin or chymotrypsin fold.

Exemplary MTSP proteins, with the protease domains indicated, are illustrated in Figures 1-3, Smaller portions thereof that retain protease activity are contemplated. The protease domains vary in size and constitution, including insertions and deletions in surface loops. They retain conserved structure, including at least one of the active site triad, primary specificity pocket, oxyanion hole and/or other features of serine protease domains of proteases. Thus, for purposes herein, the protease domain is a portion of a MTSP, as defined herein, and is homologous to a domain of other MTSPs, such as corin, enterpeptidase, human airway trypsin-like protease (HAT), MTSP1, TMPRSS2,

-17-

and TMPRSS4, which have been previously identified; it was not recognized, however, that an isolated single chain form of the protease domain could function proteolytically in *in vitro* assays. As with the larger class of enzymes of the chymotrypsin (S1) fold (see, *e.g.*, Internet accessible MEROPS data base), the MTSPs protease domains share a high degree of amino acid sequence identity. The His, Asp and Ser residues necessary for activity are present in conserved motifs. The activation site, which results in the N-terminus of second chain in the two chain forms is has a conserved motif and readily can be identified (see, *e.g.*, amino acids 801-806, SEQ ID No. 62, amino acids 406-410, SEQ ID No. 64; amino acids 186-190, SEQ ID No. 66; amino acids 161-166, SEQ ID No. 68; amino acids 255-259, SEQ ID No. 70; amino acids 190-194, SEQ ID No. 72).

As used herein, the catalytically active domain of an MTSP refers to the protease domain. Reference to the protease domain of an MTSP refers includes the single and double-chain forms of any of these proteins. The zymogen form of each protein is single chain form, which can be converted to the active two chain form by cleavage. The protease domain may also be converted to a two chain form. By active form is meant a form active *in vivo*.

15

20

25

Significantly, it is shown herein, that, at least *in vitro*, the single chain forms of the MTSPs and the catalytic domains or proteolytically active portions thereof (typically C-terminal truncations) thereof exhibit protease activity. Hence provided herein are isolated single chain forms of the protease domains of MTSPs and their use in *in vitro* drug screening assays for identification of agents that modulate the activity thereof.

As used herein an MTSP3, whenever referenced herein, includes at least one or all of or any combination of:

a polypeptide encoded by the sequence of nucleotides set forth in SEQ ID No. 3;

a polypeptide encoded by a sequence of nucleotides that

30 hybridizes under conditions of low, moderate or high stringency to the sequence of nucleotides set forth in SEQ ID No. 3;

a polypeptide that comprises the sequence of amino acids set forth as amino acids 205-437 of SEQ ID No. 4;

5

10

20

30

a polypeptide that comprises a sequence of amino acids having at least about 85% or 90% sequence identity with the sequence of amino acids set forth in SEQ ID No. 4; and/or

a splice variant of the MTSP3 set forth in SEQ ID Nos. 3 and 4.

The MTSP3 may be from any animal, particularly a mammal, and includes but are not limited to, humans, rodents, fowl, ruminants and other animals. The full length zymogen or double chain activated form is contemplated or any domain thereof, including the protease domain, which can be a double chain activated form, or a single chain form.

As used herein an MTSP4, whenever referenced herein, includes at least one or all of or any combination of:

a polypeptide encoded by the sequence of nucleotides set forth in any of SEQ ID No. 5, 7 or 9;

a polypeptide encoded by a sequence of nucleotides that hybridizes under conditions of low, moderate or high stringency to the sequence of nucleotides set forth in any of SEQ ID Nos. 5, 7 or 9;

a polypeptide that comprises the sequence of amino acids set forth in any of SEQ ID Nos. 6, 8 or 10;

a polypeptide that comprises a sequence of amino acids having at least about 85% or 90% or 95% sequence identity with the sequence of amino acids set forth in SEQ ID No. 6, 8 or 10; and/or

a splice variant of the MTSP4s set forth in SEQ ID Nos. 7-10.

The MTSP4 may be from any animal, particularly a mammal, and includes but are not limited to, humans, rodents, fowl, ruminants and other animals. The full length zymogen or double chain activated form is contemplated or any domain thereof, including the protease domain, which can be a double chain activated form, or a single chain form.

As used herein an MTSP6, whenever referenced herein, includes at least one or all of or any combination of:

a polypeptide encoded by the sequence of nucleotides set forth in any of SEQ ID No. 11;

a polypeptide encoded by a sequence of nucleotides that hybridizes under conditions of low, moderate or high stringency to the sequence of nucleotides set forth in any of SEQ ID Nos. 11;

a polypeptide that comprises the sequence of amino acids set forth in any of SEQ ID Nos. 12;

10

15

20

25

30

a polypetide that comprises a sequence of amino acids having at least about 90% or 95% or 98% sequence identity with the sequence of amino acids set forth in SEQ ID No. 12; and/or

a splice variant of the MTSP4s set forth in SEQ ID No. 12. The MTSP6 may be from any animal, particularly a mammal, and includes but are not limited to, humans, rodents, fowl, ruminants and other animals. The full length zymogen or double chain activated form is contemplated or any domain thereof, including the protease domain, which can be a double chain activated form, or a single chain form. Of particular interest herein is the MTSP6 of SEQ ID No. 12.

As used herein, a human protein is one encoded by DNA present in the genome of a human, including all allelic variants and conservative variations as long as they are not variants found in other mammals.

As used herein, a "nucleic acid encoding a protease domain or catalytically active portion of a MTSP" shall be construed as referring to a nucleic acid encoding only the recited single chain protease domain or active portion thereof, and not the other contiguous portions of the MTSP as a continuous sequence.

As used herein, a CUB domain is a motif that mediates protein-protein interactions in complement components C1r/C1s and has also been identified in various proteins involved in developmental processes.

As used herein, catalytic activity refers to the activity of the MTSP as a serine proteases. Function of the MTSP refers to its function in tumor biology, including promotion of or involvement in tumorigenesis, metastasis or carcinogenesis, and also roles in signal transduction.

As used herein, a "propeptide" or "pro sequence" is sequence of amino acids positioned at the amino terminus of a mature biologically active polypeptide. When so-positioned, the resulting polypeptide is called a zymogen. Zymogens, generally, are biologically inactive and can be converted to mature active polypeptides by catalytic or autocatalytic cleavage of the propeptide from the zymogen. A zymogen is an enzymatically inactive protein that is converted to a proteolytic enzyme by the action of an activator. Cleavage may be effected autocatalytically.

As used herein, "disease or disorder" refers to a pathological condition in an organism resulting from, e.g., infection or genetic defect, and characterized by identifiable symptoms.

As used herein, neoplasm (neoplasia) refers to abnormal new growth, and thus means the same as *tumor*, which may be benign or malignant. Unlike *hyperplasia*, neoplastic proliferation persists even in the absence of the original stimulus.

15

20

25

30

As used herein, neoplastic disease refers to any disorder involving cancer, including tumor development, growth, metastasis and progression.

As used herein, cancer refers to a general term for diseases caused by any type of malignant tumor.

As used herein, malignant, as applies to tumors, refers to primary tumors that have the capacity of *metastasis* with loss of *growth control* and *positional* control.

As used herein, an anti-cancer agent (used interchangeable with "anti-tumor or anti-neoplastic agent") refers to any agents used in the anti-cancer treatment. These include any agents, when used alone or in combination with other compounds, that can alleviate, reduce, ameliorate, prevent, or place or maintain in a state of remission of clinical symptoms or diagnostic markers associated with neoplastic disease, tumor and cancer, and can be used in methods, combinations and compositions provided herein. Non-limiting examples of anti-neoplastic agents include anti-angiogenic agents, alkylating agents, antimetabolite, certain natural products, platinum coordination complexes, anthracenediones, substituted ureas, methylhydrazine derivatives,

adrenocortical suppressants, certain hormones, antagonists and anti-cancer polysaccharides.

As used herein, a splice variant refers to a variant produced by differential processing of a primary transcript of genomic DNA that results in more than one type of mRNA. Splice variants of MTSPs are provided herein.

As used herein, angiogenesis is intended to broadly encompass the totality of processes directly or indirectly involved in the establishment and maintenance of new vasculature (neovascularization), including, but not limited to, neovascularization associated with tumors.

10

15

20

25

30

As used herein, anti-angiogenic treatment or agent refers to any therapeutic regimen and compound, when used alone or in combination with other treatment or compounds, that can alleviate, reduce, ameliorate, prevent, or place or maintain in a state of remission of clinical symptoms or diagnostic markers associated with undesired and/or uncontrolled angiogenesis. Thus, for purposes herein an anti-angiogenic agent refers to an agent that inhibits the establishment or maintenance of vasculature. Such agents include, but are not limited to, anti-tumor agents, and agents for treatments of other disorders associated with undesirable angiogenesis, such as diabetic retinopathies, restenosis, hyperproliferative disorders and others.

As used herein, non-anti-angiogenic anti-tumor agents refer to anti-tumor agents that do not act primarily by inhibiting angiogenesis.

As used herein, pro-angiogenic agents are agents that promote the establishment or maintenance of the vasculature. Such agents include agents for treating cardiovascular disorders, including heart attacks and strokes.

As used herein, undesired and/or uncontrolled angiogenesis refers to pathological angiogenesis wherein the influence of angiogenesis stimulators outweighs the influence of angiogenesis inhibitors. As used herein, deficient angiogenesis refers to pathological angiogenesis associated with disorders where there is a defect in normal angiogenesis resulting in aberrant angiogenesis or an absence or substantial reduction in angiogenesis.

As used herein, endotheliase refers to a mammalian protein, including humans, that has a transmembrane domain and is expressed on the surface of

endothelial cells and includes a protease domain, particularly an extracellular protease domain, and is preferably a serine protease. Thus, reference, for example, to endotheliase encompasses all proteins encoded by the endotheliase gene family, or an equivalent molecule obtained from any other source or that has been prepared synthetically or that exhibits the same activity. The endotheliase gene family are transmembrane proteases expressed in endothelial cells. Endotheliases are excluded from the MTSPs contemplated herein.

As used herein, the protease domain of an endotheliase refers to the polypeptide portion of the endotheliase that is the extracellular portion that exhibits protease activity. The protease domain is a polypeptide that includes at least the minimum number of amino acids, generally more than 50 or 100, required for protease activity. Protease activity may be assessed empirically, such as by testing the polypeptide for its ability to act as a protease. Assays, such as the assays described in the EXAMPLES, employing a known substrate in place of the test compounds may be used. Furthermore, since proteases, particularly serine proteases, have characteristic structures and sequences or motifs, the protease domain may be readily identified by such structure and sequence or motif.

10

15

20

As used herein, the protease domain of an MTSP protein refers to the protease domain of an MTSP that is located within or is the extracellular domain of an MTSP and exhibits serine proteolytic activity. Hence it is at least the minimal portion of the extracellular domain that exhibits proteolytic activity as assessed by standard assays *in vitro*. It refers, herein, to a single chain form heretofore thought to be inactive.

Exemplary protease domains include at least a sufficient portion of sequences of amino acids set forth as amino acids 615-855 in SEQ ID No. 2 (encoded by nucleotides 1865-2587 in SEQ ID No. 1; see also SEQ ID Nos. 49 and 50) from MTSP1, amino acids 205-437 of SEQ ID NO. 4 from MTSP3, SEQ ID No. 6, which sets forth the protease domain of MTSP4, and amino acids 217-443 of SEQ ID No. 11 from MTSP6. Also contemplated are nucleic acid molecules that encode polypeptide that has proteolytic activity in an *in vitro* proteolysis assay and that have at least 80%, 85%, 90% or 95% sequence identity with the full

length of a protease domain of an MTSP protein, or that hybridize along their full length to a nucleic acids that encode a protease domain, particularly under conditions of moderate, generally high, stringency.

5

10

15

20

25

30

For each of these protease domains, residues at the N-terminus can be critical for activity, since it has been shown that an Asp in the N-terminus of such proteases is essential for formation of the catalytically active conformation upon activation cleavage of the zymogen form of the protease. It is shown herein that the protease domain of the singles chain form of the protease is catalytically active. Hence the protease domain will require the N-terminal amino acids; the c-terminus portion may be truncated. The amount that can be removed can be determined empirically by testing the protein for protease activity in an *in vitro* assays that assesses catalytic cleavage.

Hence smaller portions of the protease domains, particularly the single chain domains, thereof that retain protease activity are contemplated. Such smaller versions will generally be C-terminal truncated versions of the protease domains. The protease domains vary in size and constitution, including insertions and deletions in surface loops. Such domains exhibit conserved structure, including at least one structural feature, such as the active site triad, primary specificity pocket, oxyanion hole and/or other features of serine protease domains of proteases. Thus, for purposes herein, the protease domain is a single chain portion of an MTSP, as defined herein, but is homologous in its structural features and retention of sequence of similarity or homology the protease domain of chymotrypsin or trypsin. Most significantly, the polypeptide will exhibit proteolytic activity as a single chain.

As used herein, by homologous means about greater than 25% nucleic acid sequence identity, preferably 25% 40%, 60%, 80%, 90% or 95%. The terms "homology" and "identity" are often used interchangeably. In general, sequences are aligned so that the highest order match is obtained (see, e.g.: Computational Molecular Biology, Lesk, A.M., ed., Oxford University Press, New York, 1988; Biocomputing: Informatics and Genome Projects, Smith, D.W., ed., Academic Press, New York, 1993; Computer Analysis of Sequence Data, Part I, Griffin, A.M., and Griffin, H.G., eds., Humana Press, New Jersey, 1994;

Sequence Analysis in Molecular Biology, von Heinje, G., Academic Press, 1987; and Sequence Analysis Primer, Gribskov, M. and Devereux, J., eds., M Stockton Press, New York, 1991; Carillo et al. (1988) SIAM J Applied Math 48:1073).

By sequence identity, the number of conserved amino acids are determined by standard alignment algorithms programs, and are used with default gap penalties established by each supplier. Substantially homologous nucleic acid molecules would hybridize typically at moderate stringency or at high stringency all along the length of the nucleic acid of interest. Also contemplated are nucleic acid molecules that contain degenerate codons in place of codons in the hybridizing nucleic acid molecule.

5

10

Whether any two nucleic acid molecules have nucleotide sequences that are at least 80%, 85%, 90%, 95%, 96%, 97%, 98% or 99% "identical" can be determined using known computer algorithms such as the "FAST A" program, using for example, the default parameters as in Pearson et al. (1988) Proc. Natl. 15 Acad. Sci. USA 85:2444 (other programs include the GCG program package (Devereux, J., et al., Nucleic Acids Research 12(I):387 (1984)), BLASTP, BLASTN, FASTA (Atschul, S.F., et al., J Molec Biol 215:403 (1990); Guide to Huge Computers, Martin J. Bishop, ed., Academic Press, San Diego, 1994, and Carillo et al. (1988) SIAM J Applied Math 48:1073). For example, the BLAST function of the National Center for Biotechnology Information database may be 20 used to determine identity. Other commercially or publicly available programs include, DNAStar "MegAlign" program (Madison, WI) and the University of Wisconsin Genetics Computer Group (UWG) "Gap" program (Madison WI)). Percent homology or identity of proteins and/or nucleic acid moleucles may be 25 determined, for example, by comparing sequence information using a GAP computer program (e.g., Needleman et al. (1970) J. Mol. Biol. 48:443, as revised by Smith and Waterman ((1981) Adv. Appl. Math. 2:482). Briefly, the GAP program defines similarity as the number of aligned symbols (i.e., nucleotides or amino acids) which are similar, divided by the total number of symbols in the shorter of the two sequences. Default parameters for the GAP 30 program may include: (1) a unary comparison matrix (containing a value of 1 for identities and 0 for non-identities) and the weighted comparison matrix of

Gribskov *et al.* (1986) *Nucl. Acids Res.* 14:6745, as described by Schwartz and Dayhoff, eds., *ATLAS OF PROTEIN SEQUENCE AND STRUCTURE*, National Biomedical Research Foundation, pp. 353-358 (1979); (2) a penalty of 3.0 for each gap and an additional 0.10 penalty for each symbol in each gap; and (3) no penalty for end gaps.

5

10

15

20

25

30

Therefore, as used herein, the term "identity" represents a comparison between a test and a reference polypeptide or polynucleotide. For example, a test polypeptide may be defined as any polypeptide that is 90% or more identical to a reference polypeptide. As used herein, the term at least "90% identical to" refers to percent identities from 90 to 99.99 relative to the reference polypeptides. Identity at a level of 90% or more is indicative of the fact that, assuming for exemplification purposes' a test and reference polynucleotide length of 100 amino acids are compared. No more than 10% (i.e., 10 out of 100) amino acids in the test polypeptide differs from that of the reference polypeptides. Similar comparisons may be made between a test and reference polynucleotides. Such differences may be represented as point mutations randomly distributed over the entire length of an amino acid sequence or they may be clustered in one or more locations of varying length up to the maximum allowable, e.g. 10/100 amino acid difference (approximately 90% identity). Differences are defined as nucleic acid or amino acid substitutions, or deletions. At level of homologies or identities above about 85-90%, the result should be independent of the program and gap parameters set; such high levels of identity readily can be assess, often without relying on software.

As used herein, primer refers to an oligonucleotide containing two or more deoxyribonucleotides or ribonucleotides, preferably more than three, from which synthesis of a primer extension product can be initiated. Experimental conditions conducive to synthesis include the presence of nucleoside triphosphates and an agent for polymerization and extension, such as DNA polymerase, and a suitable buffer, temperature and pH.

As used herein, animals include any animal, such as, but are not limited to, goats, cows, deer, sheep, rodents, pigs and humans. Non-human animals, exclude humans as the contemplated animal. The MTSPs provided herein are

-26-

from any source, animal, plant, prokaryotic and fungal. Preferred MTSPs are of animal origin, preferably mammalian origin.

As used herein, genetic therapy involves the transfer of heterologous

DNA to the certain cells, target cells, of a mammal, particularly a human, with a disorder or conditions for which such therapy is sought. The DNA is introduced into the selected target cells in a manner such that the heterologous DNA is expressed and a therapeutic product encoded thereby is produced. Alternatively, the heterologous DNA may in some manner mediate expression of DNA that encodes the therapeutic product, or it may encode a product, such as a peptide or RNA that in some manner mediates, directly or indirectly, expression of a therapeutic product. Genetic therapy may also be used to deliver nucleic acid encoding a gene product that replaces a defective gene or supplements a gene product produced by the mammal or the cell in which it is introduced. The introduced nucleic acid may encode a therapeutic compound, such as a growth factor inhibitor thereof, or a tumor necrosis factor or inhibitor thereof, such as a receptor therefor, that is not normally produced in the mammalian host or that is not produced in therapeutically effective amounts or at a therapeutically useful time. The heterologous DNA encoding the therapeutic product may be modified prior to introduction into the cells of the afflicted host in order to enhance or otherwise alter the product or expression thereof. Genetic therapy may also involve delivery of an inhibitor or repressor or other modulator of gene expression.

10

15

20

25

30

As used herein, heterologous DNA is DNA that encodes RNA and proteins that are not normally produced *in vivo* by the cell in which it is expressed or that mediates or encodes mediators that alter expression of endogenous DNA by affecting transcription, translation, or other regulatable biochemical processes. Heterologous DNA may also be referred to as foreign DNA. Any DNA that one of skill in the art would recognize or consider as heterologous or foreign to the cell in which is expressed is herein encompassed by heterologous DNA. Examples of heterologous DNA include, but are not limited to, DNA that encodes traceable marker proteins, such as a protein that confers drug resistance, DNA

that encodes therapeutically effective substances, such as anti-cancer agents,

enzymes and hormones, and DNA that encodes other types of proteins, such as antibodies. Antibodies that are encoded by heterologous DNA may be secreted or expressed on the surface of the cell in which the heterologous DNA has been introduced.

Hence, herein heterologous DNA or foreign DNA, includes a DNA molecule not present in the exact orientation and position as the counterpart DNA molecule found in the genome. It may also refer to a DNA molecule from another organism or species (*i.e.*, exogenous).

5

10

15

20

25

30

As used herein, a therapeutically effective product is a product that is encoded by heterologous nucleic acid, typically DNA, that, upon introduction of the nucleic acid into a host, a product is expressed that ameliorates or eliminates the symptoms, manifestations of an inherited or acquired disease or that cures the disease.

As used herein, recitation that a polypeptide consists essentially of the protease domain means that the only MTSP portion of the polypeptide is a protease domain or a catalytically active portion thereof. The polypeptide may optionally, and generally will, include additional non-MTSP-derived sequences of amino acids.

As used herein, cancer or tumor treatment or agent refers to any therapeutic regimen and/or compound that, when used alone or in combination with other treatments or compounds, can alleviate, reduce, ameliorate, prevent, or place or maintain in a state of remission of clinical symptoms or diagnostic markers associated with deficient angiogenesis.

As used herein, domain refers to a portion of a molecule, e.g., proteins or nucleic acids, that is structurally and/or functionally distinct from other portions of the molecule.

As used herein, protease refers to an enzyme catalyzing hydrolysis of proteins or peptides. For purposes herein, the protease domain is a single chain form of an MTSP protein. For MTSP3 and MTSP4 the protease domain also includes two chain forms.

-28-

As used herein, catalytic activity refers to the activity of the MTSP as a protease as assessed in *in vitro* proteolytic assays that detect proteolysis of a selected substrate.

5

15

20

25

30

As used herein, nucleic acids include DNA, RNA and analogs thereof, including protein nucleic acids (PNA) and mixture thereof. Nucleic acids can be single or double stranded. When referring to probes or primers, optionally labeled, with a detectable label, such as a fluorescent or radiolabel, single-stranded molecules are contemplated. Such molecules are typically of a length such that they are statistically unique and of a low copy number (typically less than 5, preferably less than 3) for probing or priming a library. Generally a probe or primer contains at least 14, 16 or 30 contiguous sequence complementary to or identical to a gene of interest. Probes and primers can be 10, 20, 30, 50, 100 or more nucleic acids long.

As used herein, nucleic acid encoding a fragment or portion of an MTSP refers to a nucleic acid encoding only the recited fragment or portion of MTSP, and not the other contiguous portions of the MTSP.

As used herein, heterologous or foreign DNA and RNA are used interchangeably and refer to DNA or RNA that does not occur naturally as part of the genome in which it is present or which is found in a location or locations in the genome that differ from that in which it occurs in nature. Heterologous nucleic acid is generally not endogenous to the cell into which it is introduced, but has been obtained from another cell or prepared synthetically. Generally, although not necessarily, such nucleic acid encodes RNA and proteins that are not normally produced by the cell in which it is expressed. Any DNA or RNA that one of skill in the art would recognize or consider as heterologous or foreign to the cell in which it is expressed is herein encompassed by heterologous DNA. Heterologous DNA and RNA may also encode RNA or proteins that mediate or alter expression of endogenous DNA by affecting transcription, translation, or other regulatable biochemical processes.

As used herein, operative linkage of heterologous DNA to regulatory and effector sequences of nucleotides, such as promoters, enhancers, transcriptional and translational stop sites, and other signal sequences refers to the relationship

-29-

between such DNA and such sequences of nucleotides. For example, operative linkage of heterologous DNA to a promoter refers to the physical relationship between the DNA and the promoter such that the transcription of such DNA is initiated from the promoter by an RNA polymerase that specifically recognizes, binds to and transcribes the DNA in reading frame.

As used herein, a sequence complementary to at least a portion of an RNA, with reference to antisense oligonucleotides, means a sequence having sufficient complementarily to be able to hybridize with the RNA, preferably under moderate or high stringency conditions, forming a stable duplex; in the case of double-stranded MTSP antisense nucleic acids, a single strand of the duplex DNA may thus be tested, or triplex formation may be assayed. The ability to hybridize depends on the degree of complementarily and the length of the antisense nucleic acid. Generally, the longer the hybridizing nucleic acid, the more base mismatches with a MTSP encoding RNA it can contain and still form a stable duplex (or triplex, as the case may be). One skilled in the art can ascertain a tolerable degree of mismatch by use of standard procedures to determine the melting point of the hybridized complex.

10

15

20

25

30

For purposes herein, conservative amino acid substitutions may be made in any of MTSPs and protease domains thereof provided that the resulting protein exhibits protease activity. Conservative amino acid substitutions, such as those set forth in Table 1, are those that do not eliminate proteolytic activity. Suitable conservative substitutions of amino acids are known to those of skill in this art and may be made generally without altering the biological activity of the resulting molecule. Those of skill in this art recognize that, in general, single amino acid substitutions in non-essential regions of a polypeptide do not substantially alter biological activity (see, e.g., Watson et al. Molecular Biology of the Gene, 4th Edition, 1987, The Bejacmin/Cummings Pub. co., p.224). Also included within the definition, is the catalytically active fragment of an MTSP, particularly a single chain protease portion. Conservative amino acid substitutions are made, for example, in accordance with those set forth in TABLE 1 as follows:

-30-

# TABLE 1

	I ADEL I		
	Original residue Ala (A)	Conservative substitution Gly; Ser, Abu	
	Arg (R)	Lys, orn	
5	Asn (N)	GIn; His	
	Cys (C)	Ser	
	Gln (Q)	Asn	
	Glu (E)	Asp	
	Gly (G)	Ala; Pro	
10	His (H)	Asn; Gln	
	lle (I)	Leu; Val; Met; Nle; Nva	
	Leu (L)	lle; Val; Met; Nle; Nv	
	Lys (K)	Arg; Gln; Glu	
	Met (M)	Leu; Tyr; IIe; NLe Val	
15	Ornitine	Lys; Arg	
	Phe (F)	Met; Leu; Tyr	
	Ser (S)	Thr	
	Thr (T)	Ser	
	Trp (W)	Tyr	
20	Tyr (Y)	Trp; Phe	
	Val (V)	lle; Leu; Met; Nle; Nv	

25

30

35

Other substitutions are also permissible and may be determined empirically or in accord with known conservative substitutions.

As used herein, Abu is 2-aminobutyric acid; Orn is ornithine.

As used herein, the amino acids, which occur in the various amino acid sequences appearing herein, are identified according to their well-known, three-letter or one-letter abbreviations. The nucleotides, which occur in the various DNA fragments, are designated with the standard single-letter designations used routinely in the art.

As used herein, a splice variant refers to a variant produced by differential processing of a primary transcript of genomic DNA that results in more than one type of mRNA.

As used herein, a probe or primer based on a nucleotide sequence disclosed herein, includes at least 10, 14, preferably at least 16 or 30 or 100 contiguous sequence of nucleotides of SEQ ID Nos. 1, 3, 5, 7, 9 or 11.

As used herein, amelioration of the symptoms of a particular disorder by administration of a particular pharmaceutical composition refers to any lessening,

whether permanent or temporary, lasting or transient that can be attributed to or associated with administration of the composition.

As used herein, antisense polynucleotides refer to synthetic sequences of nucleotide bases complementary to mRNA or the sense strand of double stranded DNA. Admixture of sense and antisense polynucleotides under appropriate conditions leads to the binding of the two molecules, or hybridization. When these polynucleotides bind to (hybridize with) mRNA, inhibition of protein synthesis (translation) occurs. When these polynucleotides bind to double stranded DNA, inhibition of RNA synthesis (transcription) occurs. The resulting inhibition of translation and/or transcription leads to an inhibition of 10 the synthesis of the protein encoded by the sense strand. Antisense nucleic acid molecule typically contain a sufficient number of nucleotides to specifically bind to a target nucleic acid, generally at least 5 contiguous nucleotides, often at least 14 or 16 or 30 contiguous nucleotides or modified nucleotides 15 complementary to the coding portion of a nucleic acid molecule that encodes a gene of interest, for example, nucleic acid encoding a single chain protease domain of an MTSP.

As used herein, an array refers to a collection of elements, such as antibodies, containing three or more members. An addressable array is one in which the members of the array are identifiable, typically by position on a solid phase support. Hence, in general the members of the array will be immobilized to discrete identifiable loci on the surface of a solid phase.

20

25

30

As used herein, antibody refers to an immunoglobulin, whether natural or partially or wholly synthetically produced, including any derivative thereof that retains the specific binding ability the antibody. Hence antibody includes any protein having a binding domain that is homologous or substantially homologous to an immunoglobulin binding domain. Antibodies include members of any immunoglobulin claims, including IgG, IgM, IgA, IgD and IgE.

As used herein, antibody fragment refers to any derivative of an antibody that is less then full length, retaining at least a portion of the full-length antibody's specific binding ability. Examples of antibody fragments include, but are not limited to, Fab, Fab', F(ab)<sub>2</sub>, single-chain Fvs (scFV), FV, dsFV diabody

and Fd fragments. The fragment can include multiple chains linked together, such as by disulfide bridges. An antibody fragment generally contains at least about 50 amino acids and typically at least 200 amino acids.

As used herein, an Fv antibody fragment is composed of one variable heavy domain ( $V_H$ ) and one variable light domain linked by noncovalent interactions.

5

10

15

20

25

30

As used herein, a dsFV refers to an Fv with an engineered intermolecular disulfide bond, which stabilizes the  $V_{H}$ - $V_{I}$  pair.

As used herein, an F(ab)<sub>2</sub> fragment is an antibody fragment that results from digestion of an immunoglobulin with pepsin at pH 4.0-4.5; it may be recombinantly produced.

As used herein, Fab fragments is an antibody fragment that results from digestion of an immunoglobulin with papain; it may be recombinantly produced.

As used herein, scFVs refer to antibody fragments that contain a variable light chain  $(V_L)$  and variable heavy chain  $(V_H)$  covalently connected by a polypeptide linker in any order. The linker is of a length such that the two variable domains are bridged without substantial interference. Preferred linkers are  $(Gly-Ser)_n$  residues with some Glu or Lys residues dispersed throughout to increase solubility.

As used herein, humanized antibodies refer to antibodies that are modified to include human sequences of amino acids so that administration to a human will not provoke an immune response. Methods for preparation of such antibodies are known. For example, the hybridoma that expresses the monoclonal antibody is altered by recombinant DNA techniques to express an antibody in which the amino acid composition of the non-variable regions is based on human antibodies. Computer programs have been designed to identify such regions.

As used herein, diabodies are dimeric scFV; diabodies typically have shorter peptide linkers than scFvs, and they preferentially dimerize.

As used herein, humanized antibodies refer to antibodies that are modified to include human sequences of amino acids so that administration to a human will not provoke an immune response. Methods for preparation of such

10

15

20

25

antibodies are known. For example, the hybridoma that expresses the monoclonal antibody is altered by recombinant DNA techniques to express an antibody in which the amino acid composition of the non-variable regions is based on human antibodies. Computer programs have been designed to identify such regions.

As used herein, production by recombinant means by using recombinant DNA methods means the use of the well known methods of molecular biology for expressing proteins encoded by cloned DNA.

As used herein the term assessing is intended to include quantitative and qualitative determination in the sense of obtaining an absolute value for the activity of an MTSP, or a domain thereof, present in the sample, and also of obtaining an index, ratio, percentage, visual or other value indicative of the level of the activity. Assessment may be direct or indirect and the chemical species actually detected need not of course be the proteolysis product itself but may for example be a derivative thereof or some further substance.

As used herein, biological activity refers to the <u>in vivo</u> activities of a compound or physiological responses that result upon <u>in vivo</u> administration of a compound, composition or other mixture. Biological activity, thus, encompasses therapeutic effects and pharmaceutical activity of such compounds, compositions and mixtures. Biological activities may be observed in <u>in vitro</u> systems designed to test or use such activities. Thus, for purposes herein the biological activity of a luciferase is its oxygenase activity whereby, upon oxidation of a substrate, light is produced.

As used herein, a combination refers to any association between two or among more items.

As used herein, a composition refers to any mixture. It may be a solution, a suspension, liquid, powder, a paste, aqueous, non-aqueous or any combination thereof.

As used herein, a conjugate refers to the compounds provided herein that include one or more MTSPs, particularly single chain protease domains thereof, and one or more targeting agents. These conjugates include those produced by recombinant means as fusion proteins, those produced by chemical means, such as by chemical coupling, through, for example, coupling to sulfhydryl groups, and those produced by any other method whereby at least one MTSP, or a domain thereof, is linked, directly or indirectly via linker(s) to a targeting agent.

As used herein, a targeting agent, is any moiety, such as a protein or effective portion thereof, that provides specific binding of the conjugate to a cell surface receptor, which, preferably, internalizes the conjugate or MTSP portion thereof. A targeting agent may also be one that promotes or facilitates, for example, affinity isolation or purification of the conjugate; attachment of the conjugate to a surface; or detection of the conjugate or complexes containing the conjugate.

10

15

20

25

30

As used herein, an antibody conjugate refers to a conjugate in which the targeting agent is an antibody.

As used herein, humanized antibodies refer to antibodies that are modified to include human sequences of amino acids so that administration to a human will not provoke an immune response. Methods for preparation of such antibodies are known. For example, the hybridoma that expresses the monoclonal antibody is altered by recombinant DNA techniques to express an antibody in which the amino acid composition of the non-variable regions is based on human antibodies. Computer programs have been designed to identify such regions.

As used herein, derivative or analog of a molecule refers to a portion derived from or a modified version of the molecule.

As used herein, fluid refers to any composition that can flow. Fluids thus encompass compositions that are in the form of semi-solids, pastes, solutions, aqueous mixtures, gels, lotions, creams and other such compositions.

As used herein, an effective amount of a compound for treating a particular disease is an amount that is sufficient to ameliorate, or in some manner reduce the symptoms associated with the disease. Such amount may

-35-

be administered as a single dosage or may be administered according to a regimen, whereby it is effective. The amount may cure the disease but, typically, is administered in order to ameliorate the symptoms of the disease. Repeated administration may be required to achieve the desired amelioration of symptoms.

5

10

15

20

25

30

As used herein equivalent, when referring to two sequences of nucleic acids means that the two sequences in question encode the same sequence of amino acids or equivalent proteins. When equivalent is used in referring to two proteins or peptides, it means that the two proteins or peptides have substantially the same amino acid sequence with only conservative amino acid substitutions (see, e.g., Table 1, above) that do not substantially alter the activity or function of the protein or peptide. When equivalent refers to a property, the property does not need to be present to the same extent [e.g., two peptides can exhibit different rates of the same type of enzymatic activity], but the activities are preferably substantially the same. Complementary, when referring to two nucleotide sequences, means that the two sequences of nucleotides are capable of hybridizing, preferably with less than 25%, more preferably with less than 15%, even more preferably with less than 5%, most preferably with no mismatches between opposed nucleotides. Preferably the two molecules will hybridize under conditions of high stringency.

As used herein, an agent that modulates the activity of a protein or expression of a gene or nucleic acid either decreases or increases or otherwise alters the activity of the protein or, in some manner up- or down-regulates or otherwise alters expression of the nucleic acid in a cell.

As used herein, inhibitor of an the activity of an MTSP encompasses any substances that prohibit or decrease production, post-translational modification(s), maturation, or membrane localization of the MTSP or any substances that interfere with or decrease the proteolytic efficacy of thereof, particular of a single chain form *in vitro*.

As used herein, a method for treating or preventing neoplastic disease means that any of the symptoms, such as the tumor, metastasis thereof, the vascularization of the tumors or other parameters by which the disease is

characterized are reduced, ameliorated, prevented, placed in a state of remission, or maintained in a state of remission. It also means that the hallmarks of neoplastic disease and metastasis may be eliminated, reduced or prevented by the treatment. Non-limiting examples of the hallmarks include uncontrolled degradation of the basement membrane and proximal extracellular matrix, migration, division, and organization of the endothelial cells into new functioning capillaries, and the persistence of such functioning capillaries.

As used herein, operatively linked or operationally associated refers to the functional relationship of DNA with regulatory and effector sequences of nucleotides, such as promoters, enhancers, transcriptional and translational stop sites, and other signal sequences. For example, operative linkage of DNA to a promoter refers to the physical and functional relationship between the DNA and the promoter such that the transcription of such DNA is initiated from the promoter by an RNA polymerase that specifically recognizes, binds to and transcribes the DNA. In order to optimize expression and/or in vitro transcription, it may be necessary to remove, add or alter 5' untranslated portions of the clones to eliminate extra, potential inappropriate alternative translation initiation (i.e., start) codons or other sequences that may interfere with or reduce expression, either at the level of transcription or translation. Alternatively, consensus ribosome binding sites (see, e.g., Kozak J. Biol. Chem. 266:19867-19870 (1991)) can be inserted immediately 5' of the start codon and may enhance expression. The desirability of (or need for) such modification may be empirically determined.

10

15

20

25

30

As used herein, pharmaceutically acceptable salts, esters or other derivatives of the conjugates include any salts, esters or derivatives that may be readily prepared by those of skill in this art using known methods for such derivatization and that produce compounds that may be administered to animals or humans without substantial toxic effects and that either are pharmaceutically active or are prodrugs.

As used herein, a prodrug is a compound that, upon in vivo administration, is metabolized or otherwise converted to the biologically, pharmaceutically or therapeutically active form of the compound. To produce a

prodrug, the pharmaceutically active compound is modified such that the active compound will be regenerated by metabolic processes. The prodrug may be designed to alter the metabolic stability or the transport characteristics of a drug, to mask side effects or toxicity, to improve the flavor of a drug or to alter other characteristics or properties of a drug. By virtue of knowledge of pharmacodynamic processes and drug metabolism in vivo, those of skill in this art, once a pharmaceutically active compound is known, can design prodrugs of the compound (see, e.g., Nogrady (1985) Medicinal Chemistry A Biochemical Approach, Oxford University Press, New York, pages 388-392).

5

10

15

20

25

30

As used herein, a drug identified by the screening methods provided herein refers to any compound that is a candidate for use as a therapeutic or as lead compound for designed a therapeutic. Such compounds can be small molecules, including small organic molecules, peptides, peptide mimetics, antisense molecules, antibodies, fragments of antibodies, recombinant antibodies and other such compound which can serve as drug candidate or lead compound.

As used herein, production by recombinant means by using recombinant DNA methods means the use of the well known methods of molecular biology for expressing proteins encoded by cloned DNA.

As used herein, a promoter region or promoter element refers to a segment of DNA or RNA that controls transcription of the DNA or RNA to which it is operatively linked. The promoter region includes specific sequences that are sufficient for RNA polymerase recognition, binding and transcription initiation. This portion of the promoter region is referred to as the promoter. In addition, the promoter region includes sequences that modulate this recognition, binding and transcription initiation activity of RNA polymerase. These sequences may be cis acting or may be responsive to trans acting factors. Promoters, depending upon the nature of the regulation, may be constitutive or regulated. Exemplary promoters contemplated for use in prokaryotes include the bacteriophage T7 and T3 promoters.

As used herein, a receptor refers to a molecule that has an affinity for a given ligand. Receptors may be naturally-occurring or synthetic molecules.

Receptors may also be referred to in the art as anti-ligands. As used herein, the

receptor and anti-ligand are interchangeable. Receptors can be used in their unaltered state or as aggregates with other species. Receptors may be attached, covalently or noncovalently, or in physical contact with, to a binding member, either directly or indirectly via a specific binding substance or linker. Examples of receptors, include, but are not limited to: antibodies, cell membrane receptors surface receptors and internalizing receptors, monoclonal antibodies and antisera reactive with specific antigenic determinants [such as on viruses, cells, or other materials], drugs, polynucleotides, nucleic acids, peptides, cofactors, lectins, sugars, polysaccharides, cells, cellular membranes, and organelles.

5

10

15

20

25

30

Examples of receptors and applications using such receptors, include but are not restricted to:

- a) enzymes: specific transport proteins or enzymes essential to survival of microorganisms, which could serve as targets for antibiotic [ligand] selection;
- b) antibodies: identification of a ligand-binding site on the antibody molecule that combines with the epitope of an antigen of interest may be investigated; determination of a sequence that mimics an antigenic epitope may lead to the development of vaccines of which the immunogen is based on one or more of such sequences or lead to the development of related diagnostic agents or compounds useful in therapeutic treatments such as for auto-immune diseases
- c) nucleic acids: identification of ligand, such as protein or RNA, binding sites;
- d) catalytic polypeptides: polymers, preferably polypeptides, that are capable of promoting a chemical reaction involving the conversion of one or more reactants to one or more products; such polypeptides generally include a binding site specific for at least one reactant or reaction intermediate and an active functionality proximate to the binding site, in which the functionality is capable of chemically modifying the bound reactant [see, e.g., U.S. Patent No. 5,215,899];
- e) hormone receptors: determination of the ligands that bind with high affinity to a receptor is useful in the development of hormone replacement therapies; for example, identification of ligands that bind to such receptors may lead to the development of drugs to control blood pressure; and

f) opiate receptors: determination of ligands that bind to the opiate receptors in the brain is useful in the development of less-addictive replacements for morphine and related drugs.

As used herein, sample refers to anything which may contain an analyte for which an analyte assay is desired. The sample may be a biological sample, such as a biological fluid or a biological tissue. Examples of biological fluids include urine, blood, plasma, serum, saliva, semen, stool, sputum, cerebral spinal fluid, tears, mucus, amniotic fluid or the like. Biological tissues are aggregate of cells, usually of a particular kind together with their intercellular substance that form one of the structural materials of a human, animal, plant, bacterial, fungal or viral structure, including connective, epithelium, muscle and nerve tissue. Examples of biological tissues also include organs, tumors, lymph nodes, arteries and individual cell(s).

As used herein: stringency of hybridization in determining percentage mismatch is as follows:

1) high stringency: 0.1 x SSPE, 0.1% SDS, 65°C

5

10

15

30

- 2) medium stringency: 0.2 x SSPE, 0.1% SDS, 50°C
- 3) low stringency: 1.0 x SSPE, 0.1% SDS, 50°C

Those of skill in this art know that the washing step selects for stable

10 hybrids and also know the ingredients of SSPE (see, e.g., Sambrook, E.F.

11 Fritsch, T. Maniatis, in: Molecular Cloning, A Laboratory Manual, Cold Spring

12 Harbor Laboratory Press (1989), vol. 3, p. B.13, see, also, numerous catalogs

13 that describe commonly used laboratory solutions). SSPE is pH 7.4 phophate
14 buffered 0.18 NaCl. Further, those of skill in the art recognize that the stability

15 of hybrids is determined by T<sub>m</sub>, which is a function of the sodium ion

16 concentration and temperature (T<sub>m</sub> = 81.5° C-16.6(log<sub>10</sub>[Na<sup>+</sup>]) + 0.41(%G+C)-600/l)), so that the only parameters in the wash conditions critical to hybrid

17 stability are sodium ion concentration in the SSPE (or SSC) and temperature.

It is understood that equivalent stringencies may be achieved using alternative buffers, salts and temperatures. By way of example and not limitation, procedures using conditions of low stringency are as follows (see also Shilo and Weinberg, *Proc. Natl. Acad. Sci. USA*, 78:6789-6792 (1981)): Filters

containing DNA are pretreated for 6 hours at 40°C in a solution containing 35% formamide, 5X SSC, 50 mM Tris-HCl (pH 7.5), 5 mM EDTA, 0.1% PVP, 0.1% FicoII, 1% BSA, and 500  $\mu$ g/ml denatured salmon sperm DNA (10X SSC is 1.5 M sodium chloride, and 0.15 M sodium citrate, adjusted to a pH of 7).

5

10

15

20

25

30

Hybridizations are carried out in the same solution with the following modifications: 0.02% PVP, 0.02% FicoII, 0.2% BSA, 100 μg/ml salmon sperm DNA, 10% (wt/vol) dextran sulfate, and 5-20 X 10<sup>6</sup> cpm <sup>32</sup>P-labeled probe is used. Filters are incubated in hybridization mixture for 18-20 hours at 40°C, and then washed for 1.5 hours at 55°C in a solution containing 2X SSC, 25 mM Tris-HCl (pH 7.4), 5 mM EDTA, and 0.1% SDS. The wash solution is replaced with fresh solution and incubated an additional 1.5 hours at 60°C. Filters are blotted dry and exposed for autoradiography. If necessary, filters are washed for a third time at 65-68°C and reexposed to film. Other conditions of low stringency which may be used are well known in the art (*e.g.*, as employed for cross-species hybridizations).

By way of example and not way of limitation, procedures using conditions of moderate stringency is provided. For example, but not limited to, procedures using such conditions of moderate stringency are as follows: Filters containing DNA are pretreated for 6 hours at 55°C in a solution containing 6X SSC, 5X Denhart's solution, 0.5% SDS and 100 μg/ml denatured salmon sperm DNA. Hybridizations are carried out in the same solution and 5-20 X 10° cpm <sup>32</sup>P-labeled probe is used. Filters are incubated in hybridization mixture for 18-20 hours at 55°C, and then washed twice for 30 minutes at 60°C in a solution containing 1X SSC and 0.1% SDS. Filters are blotted dry and exposed for autoradiography. Other conditions of moderate stringency which may be used are well-known in the art. Washing of filters is done at 37°C for 1 hour in a solution containing 2X SSC, 0.1% SDS.

By way of example and not way of limitation, procedures using conditions of high stringency are as follows: Prehybridization of filters containing DNA is carried out for 8 hours to overnight at 65°C in buffer composed of 6X SSC, 50 mM Tris-HCl (pH 7.5), 1 mM EDTA, 0.02% PVP, 0.02% Ficoll, 0.02% BSA, and 500  $\mu$ g/ml denatured salmon sperm DNA. Filters are hybridized for 48 hours

at 65°C in prehybridization mixture containing 100  $\mu$ g/ml denatured salmon sperm DNA and 5-20 X 10<sup>6</sup> cpm of <sup>32</sup>P-labeled probe. Washing of filters is done at 37°C for 1 hour in a solution containing 2X SSC, 0.01% PVP, 0.01% Ficoll, and 0.01% BSA. This is followed by a wash in 0.1X SSC at 50°C for 45 minutes before autoradiography. Other conditions of high stringency which may be used are well known in the art.

5

10

15

20

25

30

The term substantially identical or homologous or similar varies with the context as understood by those skilled in the relevant art and generally means at least 70%, preferably means at least 80%, more preferably at least 90%, and most preferably at least 95% identity.

As used herein, substantially identical to a product means sufficiently similar so that the property of interest is sufficiently unchanged so that the substantially identical product can be used in place of the product.

As used herein, substantially pure means sufficiently homogeneous to appear free of readily detectable impurities as determined by standard methods of analysis, such as thin layer chromatography (TLC), gel electrophoresis and high performance liquid chromatography (HPLC), used by those of skill in the art to assess such purity, or sufficiently pure such that further purification would not detectably alter the physical and chemical properties, such as enzymatic and biological activities, of the substance. Methods for purification of the compounds to produce substantially chemically pure compounds are known to those of skill in the art. A substantially chemically pure compound may, however, be a mixture of stereoisomers or isomers. In such instances, further purification might increase the specific activity of the compound.

As used herein, target cell refers to a cell that expresses an MTSP in vivo.

As used herein, test substance refers to a chemically defined compound (e.g., organic molecules, inorganic molecules, organic/inorganic molecules, proteins, peptides, nucleic acids, oligonucleotides, lipids, polysaccharides, saccharides, or hybrids among these molecules such as glycoproteins, etc.) or mixtures of compounds (e.g., a library of test compounds, natural extracts or culture supernatants, etc.) whose effect on an MTSP, particularly a single chain

-42-

form that includes the protease domain or a sufficient portion thereof for activity, as determined by *in vitro* method, such as the assays provided herein.

As used herein, the terms a therapeutic agent, therapeutic regimen, radioprotectant, chemotherapeutic mean conventional drugs and drug therapies, including vaccines, which are known to those skilled in the art. Radiotherapeutic agents are well known in the art.

As used herein, treatment means any manner in which the symptoms of a conditions, disorder or disease are ameliorated or otherwise beneficially altered. Treatment also encompasses any pharmaceutical use of the compositions herein.

10

15

20

25

30

As used herein, vector (or plasmid) refers to discrete elements that are used to introduce heterologous DNA into cells for either expression or replication thereof. The vectors typically remain episomal, but may be designed to effect integration of a gene or portion thereof into a chromosome of the genome. Also contemplated are vectors that are artificial chromosomes, such as yeast artificial chromosomes and mammalian artificial chromosomes. Selection and use of such vehicles are well known to those of skill in the art. An expression vector includes vectors capable of expressing DNA that is operatively linked with regulatory sequences, such as promoter regions, that are capable of effecting expression of such DNA fragments. Thus, an expression vector refers to a recombinant DNA or RNA construct, such as a plasmid, a phage, recombinant virus or other vector that, upon introduction into an appropriate host cell, results in expression of the cloned DNA. Appropriate expression vectors are well known to those of skill in the art and include those that are replicable in eukaryotic cells and/or prokaryotic cells and those that remain episomal or those which integrate into the host cell genome.

As used herein, protein binding sequence refers to a protein or peptide sequence that is capable of specific binding to other protein or peptide sequences generally, to a set of protein or peptide sequences or to a particular protein or peptide sequence.

As used herein, epitope tag refers to a short stretch of amino acid residues corresponding to an epitope to facilitate subsequent biochemical and immunological analysis of the epitope tagged protein or peptide. Epitope tagging

is achieved by appending the sequence of the epitope tag to the proteinencoding sequence in an appropriate expression vector. Epitope tagged proteins can be affinity purified using highly specific antibodies raised against the tags.

As used herein, metal binding sequence refers to a protein or peptide sequence that is capable of specific binding to metal ions generally, to a set of metal ions or to a particular metal ion.

5

15

20

25

30

As used herein, a composition refers to a any mixture. It may be a solution, a suspension, liquid, powder, a paste, aqueous, non-aqueous or any combination thereof.

As used herein, a combination refers to any association between two or among more items.

As used herein, fluid refers to any composition that can flow. Fluids thus encompass compositions that are in the form of semi-solids, pastes, solutions, aqueous mixtures, gels, lotions, creams and other such compositions.

As used herein, a cellular extract refers to a preparation or fraction which is made from a lysed or disrupted cell.

'As used herein, an agent is said to be randomly selected when the agent is chosen randomly without considering the specific sequences involved in the association of a protein alone or with its associated substrates, binding partners, etc. An example of randomly selected agents is the use a chemical library or a peptide combinatorial library, or a growth broth of an organism.

As used herein, an agent is the to be rationally selected or designed when the agent is chosen on a non-random basis which takes into account the sequence of the target site and/or its conformation in connection with the agent's action. As described in the Examples, there are proposed binding sites for serine protease and (catalytic) sites in the protein having SEQ ID NO:3 or SEQ ID NO:4. Agents can be rationally selected or rationally designed by utilizing the peptide sequences that make up these sites. For example, a rationally selected peptide agent can be a peptide whose amino acid sequence is identical to the ATP or calmodulin binding sites or domains.

For clarity of disclosure, and not by way of limitation, the detailed description is divided into the subsections that follow.

-44-

# B. MTSP PROTEINS, MUTEINS, DERIVATIVES AND ANALOGS THEREOF MTSPs

The MTSPs are a family of transmembrane serine proteases that are found in mammals and also other species that share a number of common structural features including: a proteolytic extracellular C-terminal domain; a transmembrane domain, with a hydrophobic domain near the N-terminus; a short cytoplasmic domain; and a variable length stem region containing modular domains. The proteolytic domains share sequence homology including conserved his, asp, and ser residues necessary for catalytic activity that are present in conserved motifs. The MTSPs are synthesized as zymogens, and activated to double chain forms by cleavage. It is shown herein that the single chain proteolytic domain can function *in vitro* and, hence is useful in *in vitro* assays for identifying agents that modulate the activity of members of this family. Also provided are family members designated MTSP3, MTSP4 and an MTSP6 variant.

10

15

20

25

30

The MTSP family is a target for therapeutic intervention and also some, may serve as diagnostic markers for tumor development, growth and/or progression. As discussed, the members of this family are involved in proteolytic processes that are implicated in tumor development, growth and/or progression. This implication is based upon their functions as proteolytic enzymes in

processes related to ECM degradative pathways. In addition, their levels of expression or level of activation or their apparent activity resulting from substrate levels or alterations in substrates and levels thereof differs in tumor cells and non-tumor cells in the same tissue. Hence, protocols and treatments that alter their activity, such as their proteolytic acitivities and roles in signal transduction, and/or their expression, such as by contacting them with a compound that modulates their activity and/or expression, could impact tumor development, growth and/or progression. Also, in some instances, the level of activation and/or expression may be altered in tumors, such as lung carcinoma, colon adenocarcinoma and ovarian carcinoma.

The MTSP may serve as a diagnostic marker for tumors. It is shown herein, that MTSP3 and MTSP4 and the MTSP6 variant provided herein are

-45-

expressed and/or activated in certain tumors; hence their activation or expression may serve as a diagnostic marker for tumor development, growth and/or progression. In other instances the MTSP protein can exhibit altered activity by virtue of a change in activity or expression of a co-factor therefor or a substrate therefor. In addition, in some instances, these MTSPS and/or variants thereof may be shed from cell surfaces. Detection of the shed MTSPS, particularly the extracellular domains, in body fluids, such as serum, blood, saliva, cerebral spinal fluid, synovial fluid and interstitial fluids, urine, sweat and other such fluids and secretions, may serve as a diagnostic tumor marker. In particular, detection of higher levels of such shed polypeptides in a subject compared to a subject known not to have any neoplastic disease or compared to earlier samples from the same subject, can be indicative of neoplastic disease in the subject.

5

10

15

20

25

30

Provided herein are isolated substantially pure single polypeptides that contain the protease domain of an MTSP as a single chain. The MTSPs contemplated herein are not expressed on endothelial cells, and, preferably, are expressed on tumor cells, typically at a level that differs from the level in which they are expressed in the non-tumor cell of the same type. Hence, for example, if the MTSP is expressed in an ovarian tumor cell, to be of interest herein with respect to ovarian cancer, it is expressed at the same level in non-tumor ovarian cells. MTSP protease domains include the single chain protease domains of MTSP1, MTSP3, MTSP4, MTSP6 and other such proteases, including, but are not limited to, corin, enterpeptidase, human airway trypsin-like protease (HAT), MTSP1, TMPRS2, and TMPRSS4.

Provided are the protease domains or proteins that include a portion of an MTSP that is the protease domain of any MTSP, particularly an MTSP1, MTSP3, MTSP4 and MTSP6. The protein can also include other non-MTSP sequences of amino acids, but will include the protease domain or a sufficient portion thereof to exhibit catalytic activity in any *in vitro* assay that assess such protease activity, such as any provided herein.

Also provided herein are nucleic acid molecules that encode MTSP proteins and the encoded proteins. In particular, nucleic acid molecules

-46-

encoding MTSP-3 and MTSP-4 from animals, including splice variants thereof are provided. The encoded proteins are also provided. Also provided are functional domains thereof.

In specific aspects, the MTSP protease domains, portions thereof, and muteins thereof are from or based on animal MTSPS, including, but are not limited to, rodent, such as mouse and rat; fowl, such as chicken; ruminants, such as goats, cows, deer, sheep; ovine, such as pigs; and humans.

5

10

15

20

25

30

In particular, MTSP derivatives can be made by altering their sequences by substitutions, additions or deletions that provide for functionally equivalent molecules. Due to the degeneracy of nucleotide coding sequences, other nucleic sequences which encode substantially the same amino acid sequence as a MTSP gene can be used. These include but are not limited to nucleotide sequences comprising all or portions of MTSP genes that are altered by the substitution of different codons that encode the amino acid residue within the sequence, thus producing a silent change. Likewise, the MTSP derivatives include, but are not limited to, those containing, as a primary amino acid sequence, all or part of the amino acid sequence of MTSP, including altered sequences in which functionally equivalent amino acid residues are substituted for residues within the sequence resulting in a silent change. For example, one or more amino acid residues within the sequence can be substituted by another amino acid of a similar polarity which acts as a functional equivalent, resulting in a silent alteration. Substitutes for an amino acid within the sequence may be selected from other members of the class to which the amino acid belongs. For example, the nonpolar (hydrophobic) amino acids include alanine, leucine, isoleucine, valine, proline, phenylalanine, tryptophan and methionine. The polar neutral amino acids include glycine, serine, threonine, cysteine, tyrosine, asparagine, and glutamine. The positively charged (basic) amino acids include arginine, lysine and histidine. The negatively charged (acidic) amino acids include aspartic acid and glutamic acid (see, e.g., Table 1).

In a preferred embodiment, the substantially purified MTSP protease is encoded by a nucleic acid that hybridizes to the a nucleic acid molecule containing the protease domain encoded by the nucleotide sequence set forth in

-47-

any of SEQ. ID Nos. 1, 3, 5, 7, 9 or 11 under at least moderate, generally high, stringency conditions, such that the protease domain encoding nucleic acid thereof hybridizes along its full length. In preferred embodiments the substantially purified MTSP protease is a single chain polypeptide that includes substantially the sequence of amino acids set forth in any SEQ ID Nos. 2, 4, 6, 8, 10 and 12 that encodes the protease domain. Specific sequences for the following human MTSPs and domains thereof are provided as follows: SEQ ID No. 3 MTSP3 nucleic acid sequence; SEQ ID No. 4 MTSP3 amino acid sequence; SEQ ID No. 5 MTSP4 nucleic acid encoding the protease domain; SEQ ID No. 6 MTSP4 amino acid sequence of the protease domain; SEQ ID No. 7 MTSP4-L nucleic acid sequence; SEQ ID No. 8 MTSP4-L amino acid sequence; SEQ ID No. 9 MTSP4-S nucleic acid sequence; SEQ ID No. 10 MTSP4-S amino acid sequence; SEQ ID No. 11 MTSP6 nucleic acid sequence; SEQ ID No. 12 MTSP6 amino acid sequence. SEQ ID No. 1 sets forth the nucleic acid sequence of the long form of MTSP1; SEQ ID No. 2 the encoded amino acid sequence; SEQ ID No. 49 sets forth the sequence of a protease domain of an MTSP1, and SEQ ID No. 50 the sequence of the encoded single chain protease domain thereof. Figures 1-3 depict the structural organization of the MTSP3, MTSP4 and MTSP6, respectively.

10

15

20

25

30

In particular, exemplary protease domains include at least a sufficient portion of sequences of amino acids set forth as amino acids 615-855 in SEQ ID No. 2 (encoded by nucleotides 1865-2587 in SEQ ID No. 1; see also SEQ ID Nos. 49 and 50) from MTSP1 (matriptase), amino acids 205-437 of SEQ ID No. 4 from MTSP3, SEQ ID No. 6, which sets forth the protease domain of MTSP4, and amino acids 217-443 of SEQ ID No. 11 from MTSP6. Also contemplated are nucleic acid molecules that encode a single chain MTSP protease that have proteolytic activity in an *in vitro* proteolysis assay and that have at least 60%, 70%, 80%, 85%, 90% or 95% sequence identity with the full length of a protease domain of an MTSP protein, or that hybridize along their full length to a nucleic acids that encode a protease domain, particularly under conditions of moderate, generally high, stringency. As above, the encoded polypeptides contain the protease as a single chain.

-48-

The isolated nucleic acids may include of at least 8 nucleotides of an MTSP sequence. In other embodiments, the nucleic acids may contain least 25 (continuous) nucleotides, 50 nucleotides, 100 nucleotides, 150 nucleotides, or 200 nucleotides of a MTSP sequence, or a full-length MTSP coding sequence. In another embodiment, the nucleic acids are smaller than 35, 200 or 500 nucleotides in length. Nucleic acids can be single or double stranded. Nucleic acids that hybridizes to or complementary to the foregoing sequences, in particular the inverse complement to nucleic acids that hybridizes to the foregoing sequences (i.e., the inverse complement of a nucleic acid strand has the complementary sequence running in reverse orientation to the strand so that the inverse complement would hybridize without mismatches to the nucleic acid strand; thus, for example, where the coding strand is that hybridizes to a nucleic acid with no mismatches between the coding strand and the that hybridizes strand, then the inverse complement of the that hybridizes strand is identical to the coding strand) are also provided. In specific aspects, nucleic acids are provided that include a sequence complementary to (specifically are the inverse complement of) at least 10, 25, 50, 100, or 200 nucleotides or the entire coding region of an MTSP encoding nucleic acid, particularly the protease domain thereof. For MTSP3 and MTSP4 the full-length protein or domain or active fragment thereof.

10

15

20

25

30

For each of the nucleic acid molecules, the nucleic acid can be DNA or RNA or PNA or other nucleic acid analogs or may include non-natural nucleotide bases.

Also provided are isolated nucleic acid molecules that include a sequence of nucleotides complementary to the nucleotide sequence encoding an MTSP.

Probes and primers derived from the nucleic acid molecules are provided, Such probes and primers contain at least 8, 14, 16, 30, 100 or more contiguous nucleotides with identity to contiguous nucleotides of an MTSP, including, but are not limited to, MTSP1, MTSP3, MTSP4 and MTSP6. The probes and primers are optionally labelled with a detectable label, such as a radiolabel or a fluorescent tag, or can be mass differentiated for detection by mass spectrometry or other means.

WO 01/57194

10

15

20

25

30

PCT/US01/03471

Plasmids and vectors containing the nucleic acid molecules are also provided. Cells containing the vectors, including cells that express the encoded proteins are provided. The cell can be a bacterial cell, a yeast cell, a fungal cell, a plant cell, an insect cell or an animal cell. Methods for producing an MTSP or single chain form of the protease domain thereof by, for example, growing the cell under conditions whereby the encoded MTSP is expressed by the cell, and recovering the expressed protein, are provided herein. As noted, for MTSP3 and MTSP4, the full-length zymogens and activated proteins and activated (two strand) protease and single chain protease domains are provided.

Except for the MTSP proteins (MTSP3 and MTSP4) heretofore unidentified and provided herein, the isolated polypeptides contain the MTSP protease domain or a catalytically active portion thereof and, generally, do not contain additional MTSP. Hence isolated, substantially pure proteases, protease domains or catalytically active portion thereof in single chain form of MTSPs are provided. The protease domains may be included in a longer protein, but such longer protein is not the MTSP zymogen.

Thus, MTSP3 and MTSP4 proteins are provided. For these proteins, the domains, fragments, derivatives or analogs that are functionally active, *i.e.*, capable of exhibiting one or more functional activities associated with the MTSP protein, *e.g.*, serine protease activity, immunogenicity and antigenicity, are provided. As discussed above, the protease domains thereof are also provided. For MTSP3 and MTSP4, the zymogens and activated forms, and also, the single chain and double chain, activated protease domains are provided.

Also provided are nucleic acid molecules that hybridize to the abovenoted sequences of nucleotides encoding MTSP3 and MTSP4 (SEQ ID Nos. 3, 5,
7 and 9) at least at low stringency, more preferably at moderate stringency, and
most preferably at high stringency, and that encode the protease domain and/or
the full length protein or other domains of an MTSP family member, such as
MTSP3, MTSP4, MTSP6 or a splice variant or allelic variant thereof, or MTSP6
or a splice variant or allelic variant thereof. Preferably the molecules hybridize
under such conditions along their full length for at least one domain and encode
at least one domain, such as the protease or extracellular domain, of the

WO 01/57194

polypeptide. In particular, such nucleic acid molecules include any isolated nucleic fragment that encodes at least one domain of a membrane serine protease, that (1) contains a sequence of nucleotides that encodes the protease or a domain thereof, and (2) is selected from among:

5

10

15

- (a) a sequence of nucleotides that encodes the protease or a domain thereof includes a sequence of nucleotides set forth above;
- (b) a sequence of nucleotides that encodes such portion or the full length protease and hybridizes under conditions of high stringency, preferably to nucleic acid that is complementary to a mRNA transcript present in a mammalian cell that encodes such protein or fragment thereof;
- a sequence of nucleotides that encodes a transmembrane protease or domain thereof that includes a sequence of amino acids
   encoded by such portion or the full length open reading frame; and
- (d) a sequence of nucleotides that encodes the transmembrane protease that includes a sequence of amino acids encoded by a sequence of nucleotides that encodes such subunit and hybridizes under conditions of high stringency to DNA that is complementary to the mRNA transcript.

## 20

## Exemplary MTSPs

The above discussion provides an overview and some details of the exemplified MTSPs. The following discussion provides additional details (see, also, EXAMPLES).

## MTSP1 (matriptase)

25 Matriptase is a trypsin-like serine protease with broad spectrum cleavage activity and two potential regulatory modules. It was named "matriptase" because its ability to degrade the extra-cellular matrix and its trypsin-like activity. When isolated from breast cancer cells (or T-47D cell conditioned medium), matriptase has been reported to be primarily in an uncomplexed form.

30 Matriptase has been isolated from human milk; when isolated from human milk, matriptase was reported to be in one of two complexed forms, 95 kDa (the predominant form) and 110 kDa; uncomplexed matriptase was not detected.

-51-

(Liu, et al., J. Biol. Chem. 274(26):18237-18242 (1999).) It has been proposed that matriptase exists as an uncomplexed protease when in its active state. In breast milk, matriptase has been reported to exist in complex with a fragment of hepatocyte growth factor inhibitor-1 (HAI-1), a Kuntz-type serine protease inhibitor having activity against trypsin-like serine proteases.

5

10

15

20

25

Ecotin and Ecotin M84R/M85R are macromolecular inhibitors of serine proteases of the chymotrypsin fold and inhibit ductal branching, morphogenesis and differentiation of the explanted ductal prostate. PC-3 is a cell line derived from prostate cancer epithelial cells. Ecotin and M84R/M85R ecotin were found to decrease tumor size and metastasis in PC-3 implanted nude mice.

Matriptase has been isolated and its encoding nucleic acids cloned from T-47D human breast cancer cell-conditioned medium (Lin *et al.* (1999) *J. Biol. Chem. 274*:18231-18236). Upon analysis of the cDNA, it was determined that the full length protease has 683 amino acids and contains three main structural regions: a serine protease domain near the carboxyl-terminal region, four tandem low-density lipoprotein receptor domains, and two tandem complement subcomponents C1r and C1s.

Studies to identify additional serine proteases made by cancer cells were done using PC-3 cells. A serine protease termed "MT-SP1", reported to be a transmembrane protease was cloned (Takeuchi *et al.* (1999) *Proc. Natl. Acad. Sci. U.S.A. 96*:11054-11061). It was subsequently found the originally identified matriptase sequence is included in the translated sequence of the cDNA that encodes MT-SP1. The matriptase cDNA was reported to be a partial MT-SP1 cDNA and to lack 516 of the coding nucleotides (Takeuchi, *et al.*, *J. Biol. Chem 275*:26333-26342 (2000).) Since the reported matriptase encoding cDNA sequence encoded a possible initiating methionine, it was proposed that alternative splicing could yield a protein lacking the N-terminal region of MTSP1.

Matriptase and MT-SP1 demonstrate trypsin-like protease activity and are

Type II transmembrane proteins with a common extracellular protease domain.

Studies of substrate specificity of MT-SP1 reveal that protease-activated receptor 2 (PAR2) and single-chain urokinase-type plasminogen activator (sc-

uPA) are macromolecular substrates of MT-SP1. PAR2 is functions in inflammation, cytoprotection and/or cell adhesion, while sc-uPa is functions in tumor cell invasion and metastasis.

5

10

15

20

25

30

An exemplary nucleotide sequences encoding a human MTSP1 is set forth in SEQ ID Nos 1 and 2 (see also SEQ ID Nos. 49 and 50 for the protease domain thereof). As previously noted SEQ ID No. 1 sets for an MTSP1-encoding nucleic acid sequence. This sequence is the longer version and includes the protease domain, which is common to both variants Nucleic acids encoding the MTSP that hybridizes to the nucleotide sequence set forth in SEQ ID No. 1 can be obtained by any method known in the art, e.g, by PCR amplification using synthetic primers that hybridizes to the 3' and 5' ends of the sequence and/or by cloning from a cDNA or genomic library using a PCR amplification product or an oligonucleotide specific for the gene sequence (e.g., as described in Section C herein). Homologs (e.g., nucleic acids of the above-listed genes of species other than human) or other related sequences (e.g., paralogs) and muteins can be obtained by low, moderate or high stringency hybridization with all or a portion of the particular sequence provided as a probe using methods well known in the art for nucleic acid hybridization and cloning.

Isolated single chain protease domains of MTSP1 proteins from animals are provided herein. As shown herein, the single chain protease domain is catalytically active and can be used in a variety of drug screening assays, particularly *in vitro* proteolytic assays. Exemplary MTSP protease domains are set forth as the amino acids (615-855 of SEQ ID No. 2) encoded by nucleotides 1865-2587 of SEQ ID No. 1 (see, also, SEQ ID Nos. 49 and 50). The MTSP1 single chain protease domain is catalytically active

Muteins of the MTSP1 proteins are provided. In the activated double chain molecule, residue 731 forms a disulfide bond with the Cys at residue 604. In the single chain form, the residue at 731 in the protease domain is free. Muteins in which Cys residues, particularly the free Cys residue (amino acid 731 in SEQ ID No. 2) in the single chain protease domain are provided. Other muteins in which conservative amino acids replacements are effected and that retain proteolytic activity as a single chain are also provided. Such changes may

-53-

be systematically introduced and tested for activity in *in vitro* assays, such as those provided herein.

#### MTSP3

5

10

15

20

25

30

In a specific embodiment, a nucleic acid that encodes a MTSP, designated MTSP3 is provided. In particular, the nucleic acid includes an open reading frame within the following sequence of nucleotides set forth in SEQ ID No. 3. In particular the protein is encoded by the open reading frame that begins at nucleotide 261 and ends at 1574.

Also provided are nucleic acid molecules that hybridize under conditions of at least low stringency, preferably moderate stringency, more preferably high stringency to the following sequence of nucleic acids (SEQ ID No. 3), particularly to the open reading frame encompassed by nucleotides that encode a single protease domain thereof, or any domain of MTSP3

Also included are substantially purified MTSP3 zymogen, activated double chains, single chain protease domains and double chain protease domains. These are encoded by a nucleic acid that includes sequence encoding a protease domain that exhibits proteolytic activity and that hybridizes to a nucleic acid molecule having a nucleotide sequence set forth in SEQ ID No. 3, typically under moderate, generally under high stringency, conditions and most preferably along the full length of the protease domain. Splice variants are also contemplated herein.

In a preferred embodiment, the isolated nucleic acid fragment hybridizes to the nucleic acid having the nucleotide sequence set forth in SEQ ID No: 3 under high stringency conditions, and preferably comprises the sequence of nucleotides set forth in any of SEQ ID Nos. 3 or comprises a portion thereof that encodes a transmembrane domain and may additionally include a LDLR domain, a scavenger-receptor cysteine rich (SRCR) domain and a serine protease catalytic domain or any other identified domain (see FIGURES) or comprises nucleic acid molecule that encodes the protein encoded by SEQ ID NO. 4.

The isolated nucleic acid fragment is DNA, including genomic or cDNA, or is RNA, or can include other components, such as protein nucleic acid. The isolated nucleic acid may include additional components, such as heterologous or

native promoters, and other transcriptional and translational regulatory sequences, these genes may be linked to other genes, such as reporter genes or other indicator genes or genes that encode indicators.

Also provided is an isolated nucleic acid molecule that includes the sequence of molecules that is complementary to the nucleotide sequence encoding the MTSP or the portion thereof.

5

10

15

20

25

30

Also provided are fragments thereof that can be used as probes or primers and that contain at least about 10 nucleotides, more preferably 14 nucleotides, more preferably at least about 16 nucleotides, most preferably at least about 30 nucleotides.

Hence provided herein are polypeptides that are encoded by such nucleic acid molecules. Included among those polypeptides are the MTSP3 protease domain or a polypeptide with conservative amino acid changes such that the specificity and protease activity remains substantially unchange. In particular, a substantially purified mammalian MTSP protein is provided that has a transmembrane domain and may additionally include a CUB domain, one or more of an LDLR domain(s), a scavenger-receptor cysteine rich (SRCR) domain and a serine protease catalytic domain is provided.

Also provided is a substantially purified protein comprising a sequence of amino acids that has at least 60%, more preferably at least about 90%, most preferably at least about 95%, identity to the MTSP3, wherein the percentage identity is determined using standard algorithms and gap penalties that maximize the percentage identity. The human MTSP3 protein is most preferred, although other mammalian MTSP3 proteins are contemplated.

Muteins of MTSP3, particularly those in which Cys residues, such as the Cys310 in the single chain protease domain, is replaced with another amino acid that does not eliminate the activity, are provided.

# MTSP4

Among the proteins provided herein is MTSP4. MTSP4 is highly expressed in the liver, and is expressed in substantially lower levels in other tissues (see, EXAMPLES). It is also expressed in non-liver-derived tumors (see EXAMPLES), including Burkitt's lymphoma, colorectal adenocarcinoma

-55-

(SW480), lung carcinoma (A549), and in leukemic cells, indicating a role in one or more of tumor progression, tumor invasion, tumor growth and tumor metastases.

Also provided are nucleic acid molecules that hybridize under conditions of at least low stringency, preferably moderate stringency, more preferably high stringency to the sequence of nucleic acids set forth in SEQ ID Nos. 5, 7 or 9), particularly to the open reading frame encompassed by nucleotides that encode a single protease domain thereof, or any domain of an MTSP4.

5

10

15

20

25

30

Also included are substantially purified MTSP4 zymogens, activated double chains, single chain protease domains and double chain protease domains. These are encoded by a nucleic acid that includes sequence encoding a protease domain that exhibits proteolytic activity and that hybridizes to a nucleic acid molecule having a nucleotide sequence set forth in SEQ ID Nos. 5, 7 and 9, typically under moderate, generally under high stringency, conditions and most preferably along the full length of the protease domain.

In a preferred embodiment, the isolated nucleic acid fragment hybridizes to the nucleic acid having the nucleotide sequence set forth in SEQ ID No: 5, 7 or 9 under high stringency conditions, and preferably comprises the sequence of nucleotides set forth in any of SEQ ID Nos. 5, 7 or 9 comprises a portion thereof that encodes a transmembrane domain and may additionally include a LDLR domain, a scavenger-receptor cysteine rich (SRCR) domain and a serine protease catalytic domain or any other identified domain (see FIGURES) or comprises nucleic acid molecule that encodes the protein encoded by SEQ ID NO. 6, 9 or 10..

The isolated nucleic acid fragment is DNA, including genomic or cDNA, or is RNA, or can include other components, such as protein nucleic acid. The isolated nucleic acid may include additional components, such as heterologous or native promoters, and other transcriptional and translational regulatory sequences, these genes may be linked to other genes, such as reporter genes or other indicator genes or genes that encode indicators.

-56-

Also provided is an isolated nucleic acid molecule that includes the sequence of molecules that is complementary to the nucleotide sequence encoding and MTSP4 or the portion thereof.

5

10

15

20

25

30

Also provided are fragments thereof that can be used as probes or primers and that contain at least about 10 nucleotides, more preferably 14 nucleotides, more preferably at least about 16 nucleotides, most preferably at least about 30 nucleotides.

In particular nucleic acid molecules encoding two forms of MTSP4 are provide. The encoded proteins are multi-domain, type II membrane-type serine proteases and include a transmembrane domain at the N terminus followed by a CUB domain, 3 LDLR domains and a trypsin-like serine protease domain at the C terminus. The difference between the two forms, which are splice variants, is the absence in MTSP4-S of a 432-bp nucleotide sequence between the transmembrane and the CUB domains (see FIGURE 2; see, also SEQ ID Nos. 5-10).

Also provided is a nucleic acid that encodes the extracellular protease domain of an MTSP4 is provided. Both forms of MTSP4 exemplified herein include a protease domain in common (see SEQ ID Nos. 5 and 6).

In particular, the extracellular protease domain of the MTSP4 proteins is encoded by the open reading frame that begins at nucleotide 1 and ends at 708 (TGA) (SEQ ID No. 5. This open reading frame encodes a portion of the MSTP4 protein and includes the protease domain. Full length MSTP4 proteins (SEQ ID Nos. 7 and 9) include the above domain. The extracellular protease domain, as a single chain, and also an activated double chain, exhibit protease activity. The disulfide bonds that form that two chain form of MTSP forms are likely between Cys415 and Cys535 for MTSP4-S, and between Cys559 and Cys679 for MTSP4-L.

For use of the single chain protease domain thereof, it is of interest to replace the free Cys (i.e. Cys535 (Cys679)) in the protease domain with another amino acid, such as any amino acid that does not alter the function (such change is likely to be any amino acid). Thus, muteins of MTSP4, particularly

-57-

those in which Cys residues, such as the Cys535 and Cys679 in the single chain protease domains of MTSP4-S and MTSP4-L, respectively, are provided.

## MTSP6

20

25

30

Nucleic acid and the encoded MTSP6 protein of an exemplary MTSP6 are
also provided. The respective sequences are set forth in SEQ ID Nos. 11 and
12. The MTSP6 DNA and protein sequences were analyzed using DNA Strider
(version 1.2). The ORF encoding the MTSP6 variant provided herein is
composed of 1,362 bp, which translate into a 453-amino acid protein. MTSP6
is a multi-domain, type-II membrane-type serine protease containing a
transmembrane domain (amino acids 48-68) at the N-terminus followed by a
LDLRa domain (LDL receptor domain class a) (amino acids 72-108), a SR domain
(Scavenger receptor Cys-rich domain)(amino acids 109-205), and a trypsin-like
serine protease domain (amino acids 216-443) (see FIGURE 3). Muteins of
MTSP6, particularly those in which Cys residues, such as the Cys324 in the
single chain protease domain of MTSP6 are provided.

International PCT application No. WO 00/52044 describes MTSPs that resemble the MTSP6 provided herein. The polypeptide provided therein differs at single amino acid positions, such as 90 in SEQ ID No. 12 (Ala is replaced with a Thr), and significantly from the instant MTSP6 in that ten amino acids (amino acid nos. 46-55 in SEQ ID No. 12) are replaced with the eleven amino acids: phe glu val phe ser gln ser ser ser leu gly (SEQ ID No. 59) resulting in a protein that is one 454 amino acids long.

There are a few other amino acid sequence differences and a number of nucleic acid sequence differences. Significantly, there are substantial differences in the protease domain at amino acids 368-394 (368 ICNHRDVYGGIISPSMLCAGYLTGGVD----- 394; SEQ ID No. 12) are replaced at position 369-396 with animo acids:

369 DLQPQ----GRVRWHHLPLHALRGLPDGWRWN 396, where the differences from 368-394 (Seq ID No. 12) are indicated.

In addition, a second C-terminus truncated variant with an altered protease domain is identified in the PCT application. The variant is the same as the 454 variant through amino acid 261 thereof (corresponding to 160 of SEQ

-58-

ID No. 12 herein), followed by 33 amino acids (see SEQ ID No. 60 herein) that differ by virtue of a frame shift.

# C. Tumor specificity and tissue expression profiles

Each MTSP has a characteristic tissue expression profile; the MTSPs in particular, although not exclusively expressed or activated in tumors, exhibit characteristic tumor tissue expression or activation profiles. In some instances, MTSPs may have different activity in a tumor cell from a non-tumor cell by virtue of a change in a substrate or cofactor thereof or other factor that would alter the apparent functional activity of the MTSP. Hence each can serve as a diagnostic marker for particular tumors, by virtue of a level of activity and/or expression or function in a subject (i.e. a mammal, particularly a human) with neoplastic disease, compared to a subject or subjects that do not have the neoplastic disease. In addition, detection of activity (and/or expression) in a particular tissue can be indicative of neoplastic disease. Shed MTSPs in body fluids can be indicative of neoplastic disease. Also, by virtue of the activity and/or expression profiles of each, they can serve as therapeutic targets, such as by administration of modulators of the activity thereof, or, as by administration of a prodrug specifically activated by one of the MTSPs.

# Tissue expression profiles

20 MTSP3

5

10

15

25

30

The MTSP3 transcript was detected in lung carcinoma (LX-1), colon adenocarcinoma (CX-1), colon adenocarcinoma (GI-112) and ovarian carcinoma (GI-102). No apparent signal was detected in another form of lung carcinoma (GI-117), breast carcinoma (GI-101), pancreatic adenocarcinoma (GI-103) and prostatic adenocarcinoma (PC3).

MTSP1 is expressed in breast cancers.

#### MTSP4

The MTSP4 transcript, a DNA fragment encoding part of the LDL receptor domain and the protease domain was used to probe an RNA blot composed of 76 different human tissues (catalog number 7775-1; human multiple tissue expression (MTE) array; CLONTECH). As in the northern analysis of gel blot, a very strong signal was observed in the liver. Signals in other tissues were

-59-

observed in (decreasing signal level): fetal liver > heart = kidney = adrenal gland = testis = fetal heart and kidney = skeletal muscle = bladder = placenta > brain = spinal cord = colon = stomach = spleen = lymph node = bone marrow = trachea = uterus = pancreas = salivary gland = mammary gland = lung. MTSP4 is also expressed less abundantly in several tumor cell lines including HeLa S3 = leukemia K-562 = Burkitt's lymphomas (Raji and Daudi) = colorectal adenocarcinoma (SW480) > lung carcinoma (A549) = leukemia MOLT-4 = leukemia HL-60. PCR of the MTSP4 transcript from cDNA libraries made from several human primary tumors xenografted in nude mice (human 10 tumor multiple tissue cDNA panel, catalog number K1522-1, CLONTECH) was performed using MTSP4-specific primers. The MTSP4 transcript was detected in breast carcinoma (GI-101), lung carcinoma (LX-1), colon adenocarcinoma (Gl-112) and pancreatic adenocarcinoma (Gl-103). No apparent signal was detected in another form of lung carcinoma (GI-117), colon adenocarcinoma (CX-1), ovarian carcinoma (GI-102) and prostatic adenocarcinoma (PC3). The 15 MTSP4 transcript was also detected in LNCaP and PC-3 prostate cancer cell lines as well as in HT-1080 human fibrosarcoma cell line.

# Gene expression profile of MTSP6 in normal and tumor tissues

To obtain information regarding the gene expression profile of the MTSP6 transcript, a 495 bp DNA fragment obtained from PCR reaction with primers Ch17-NSP-3 and NSP-4AS was used to probe an RNA blot composed of 76 different human tissues (catalog number 7775-1; human multiple tissue expression (MTE) array; CLONTECH). The strongest signal was observed in duodenum. Signal in other tissues were observed in (decreased signal level):

Stomach > trachea = mammary gland = thyroid gland = salivary gland = pituitary gland = pancreas > kidney > lung > jejunum = ileum = ilocecum = appendix = fetal kidney > fetal lung. Very weak signals can also be detected in several other tissues.

MTSP6 is also expressed in several tumor cell lines including HeLa S3 > colorectal adenocarcinoma (SW480) > leukemia MOLT-4 > leukemia K-562.

PCR analysis of the MTSP6 transcript from cDNA libraries made from several

`-60-

human primary tumors xenografted in nude mice (human tumor multiple tissue cDNA panel, catalog number K1522-1, CLONTECH) was performed using MTSP6-specific primers (Ch17-NSP-3 and Ch17-NSP2AS). The MTSP6 transcript was strongly detected in lung carcinoma (LX-1), moderately detected in pancreatic adenocarcinoma (GI-103), weakly detected in ovarian carcinoma (GI-102); and very weakly detected in colon adenocarcinoma (GI-112 and CX-1), breast carcinoma (GI-101), lung carcinoma (GI-117) and prostatic adenocarcinoma (PC3). The MTSP6 transcript was also detected in breast cancer cell line MDA-MB-231, prostate cancer cell line PC-3, but not in HT-1080 human fibrosarcoma cell line. MTSP6 is also expressed in mammary gland carcinoma cDNA (Clontech). MTSP6 is also over expressed in ovarian tumor cells.

# D. Identification and isolation of MTSP protein genes

10

15

20

25

30

The MTSP proteins, or domains thereof, can be obtained by methods well known in the art for protein purification and recombinant protein expression. Any method known to those of skill in the art for identification of nucleic acids that encode desired genes may be used. Any method available in the art can be used to obtain a full length (*i.e.*, encompassing the entire coding region) cDNA or genomic DNA clone encoding an MTSP protein. In particular, the polymerase chain reaction (PCR) can be used to amplify a sequence identified as being differentially expressed in normal and tumor cells or tissues, *e.g.*, nucleic acids encoding an MTSP protein (SEQ. NOs: 1-12), in a genomic or cDNA library. Oligonucleotide primers that hybridize to sequences at the 3' and 5' termini of the identified sequences can be used as primers to amplify by PCR sequences from a nucleic acid sample (RNA or DNA), preferably a cDNA library, from an appropriate source (*e.g.*, tumor or cancer tissue).

PCR can be carried out, *e.g.*, by use of a Perkin-Elmer Cetus thermal cycler and Taq polymerase (Gene Amp<sup>™</sup>). The DNA being amplified can include mRNA or cDNA or genomic DNA from any eukaryotic species. One can choose to synthesize several different degenerate primers, for use in the PCR reactions. It is also possible to vary the stringency of hybridization conditions used in priming the PCR reactions, to amplify nucleic acid homologs (*e.g.*, to obtain

5

10

15

20

25

30

-61-

MTSP protein sequences from species other than humans or to obtain human sequences with homology to MTSP protein) by allowing for greater or lesser degrees of nucleotide sequence similarity between the known nucleotide sequence and the nucleic acid homolog being isolated. For cross species hybridization, low stringency conditions are preferred. For same species hybridization, moderately stringent conditions are preferred. After successful amplification of the nucleic acid containing all or a portion of the identified MTSP protein sequence or of a nucleic acid encoding all or a portion of an MTSP protein homolog, that segment may be molecularly cloned and sequenced, and used as a probe to isolate a complete cDNA or genomic clone. This, in turn, will permit the determination of the gene's complete nucleotide sequence, the analysis of its expression, and the production of its protein product for functional analysis. Once the nucleotide sequence is determined, an open reading frame encoding the MTSP protein gene protein product can be determined by any method well known in the art for determining open reading frames, for example, using publicly available computer programs for nucleotide sequence analysis. Once an open reading frame is defined, it is routine to determine the amino acid sequence of the protein encoded by the open reading frame. In this way, the nucleotide sequences of the entire MTSP protein genes as well as the amino acid sequences of MTSP protein proteins and analogs may be identified.

Any eukaryotic cell potentially can serve as the nucleic acid source for the molecular cloning of the MTSP protein gene. The nucleic acids can be isolated from vertebrate, mammalian, human, porcine, bovine, feline, avian, equine, canine, as well as additional primate sources, insects, plants, etc. The DNA may be obtained by standard procedures known in the art from cloned DNA (e.g., a DNA "library"), by chemical synthesis, by cDNA cloning, or by the cloning of genomic DNA, or fragments thereof, purified from the desired cell (see, for example, Sambrook et al., 1989, Molecular Cloning, A Laboratory Manual, 2d Ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York; Glover, D.M. (ed.), 1985, DNA Cloning: A Practical Approach, MRL Press, Ltd., Oxford, U.K. Vol. I, II). Clones derived from genomic DNA may contain regulatory and intron DNA regions in addition to coding regions; clones derived

-62-

from cDNA will contain only exon sequences. Whatever the source, the gene should be molecularly cloned into a suitable vector for propagation of the gene.

In the molecular cloning of the gene from genomic DNA, DNA fragments are generated, some of which will encode the desired gene. The DNA may be cleaved at specific sites using various restriction enzymes. Alternatively, one may use DNAse in the presence of manganese to fragment the DNA, or the DNA can be physically sheared, for example, by sonication. The linear DNA fragments can then be separated according to size by standard techniques, including but not limited to, agarose and polyacrylamide gel electrophoresis and column chromatography.

10

15

20

25

30

Once the DNA fragments are generated, identification of the specific DNA fragment containing the desired gene may be accomplished in a number of ways. For example, a portion of the MTSP protein (of any species) gene (e.g., a PCR amplification product obtained as described above or an oligonucleotide having a sequence of a portion of the known nucleotide sequence) or its specific RNA, or a fragment thereof be purified and labeled, and the generated DNA fragments may be screened by nucleic acid hybridization to the labeled probe (Benton and Davis, Science 196:180 (1977); Grunstein and Hogness, Proc. Natl. Acad. Sci. U.S.A. 72:3961 (1975)). Those DNA fragments with substantial homology to the probe will hybridize. It is also possible to identify the appropriate fragment by restriction enzyme digestion(s) and comparison of fragment sizes with those expected according to a known restriction map if such is available or by DNA sequence analysis and comparison to the known nucleotide sequence of MTSP protein. Further selection can be carried out on the basis of the properties of the gene. Alternatively, the presence of the gene may be detected by assays based on the physical, chemical, or immunological properties of its expressed product. For example, cDNA clones, or DNA clones which hybrid-select the proper mRNA, can be selected which produce a protein that, e.g., has similar or identical electrophoretic migration, isolectric focusing behavior, proteolytic digestion maps, antigenic properties, serine protease activity. If an anti-MTSP protein antibody is available, the protein may be identified by binding of labeled antibody to the putatively MTSP protein

-63-

synthesizing clones, in an ELISA (enzyme-linked immunosorbent assay)-type procedure.

Alternatives to isolating the MTSP protein genomic DNA include, but are not limited to, chemically synthesizing the gene sequence from a known sequence or making cDNA to the mRNA that encodes the MTSP protein. For example, RNA for cDNA cloning of the MTSP protein gene can be isolated from cells expressing the protein. The identified and isolated nucleic acids can then be inserted into an appropriate cloning vector. A large number of vector-host systems known in the art may be used. Possible vectors include, but are not limited to, plasmids or modified viruses, but the vector system must be compatible with the host cell used. Such vectors include, but are not limited to, bacteriophages such as lambda derivatives, or plasmids such as pBR322 or pUC plasmid derivatives or the Bluescript vector (Stratagene, La Jolla, CA). The insertion into a cloning vector can, for example, be accomplished by ligating the DNA fragment into a cloning vector which has complementary cohesive termini. I the complementary restriction sites used to fragment the DNA are not present in the cloning vector, the ends of the DNA molecules may be enzymatically modified. Alternatively, any site desired may be produced by ligating nucleotide sequences (linkers) onto the DNA termini; these ligated linkers may comprise specific chemically synthesized oligonucleotides encoding restriction endonuclease recognition sequences. In an alternative method, the cleaved vector and MTSP protein gene may be modified by homopolymeric tailing. Recombinant molecules can be introduced into host cells via transformation, transfection, infection, electroporation, etc., so that many copies of the gene sequence are generated.

10

15

20

25

30

In an alternative method, the desired gene may be identified and isolated after insertion into a suitable cloning vector in a "shot gun" approach. Enrichment for the desired gene, for example, by size fractionization, can be done before insertion into the cloning vector.

In specific embodiments, transformation of host cells with recombinant DNA molecules that incorporate the isolated MTSP protein gene, cDNA, or synthesized DNA sequence enables generation of multiple copies of the gene.

-64-

Thus, the gene may be obtained in large quantities by growing transformants, isolating the recombinant DNA molecules from the transformants and, when necessary, retrieving the inserted gene from the isolated recombinant DNA.

E. Vectors, plasmids and cells that contain nucleic acids encoding an MTSP protein or protease domain thereof and expression of MTSP proteins

#### Vectors and cells

5

10

15

20

25

30

For recombinant expression of one or more of the MTSP proteins, the nucleic acid containing all or a portion of the nucleotide sequence encoding the MTSP protein can be inserted into an appropriate expression vector, *i.e.*, a vector that contains the necessary elements for the transcription and translation of the inserted protein coding sequence. The necessary transcriptional and translational signals can also be supplied by the native promoter for MTSP genes, and/or their flanking regions.

Also provided are vectors that contain nucleic acid encoding the MTSPs. Cells containing the vectors are also provided. The cells include eukaryotic and prokaryotic cells, and the vectors are any suitable for use therein.

Prokaryotic and eukaryotic cells, including endothelial cells, containing the vectors are provided. Such cells include bacterial cells, yeast cells, fungal cells. plant cells, insect cells and animal cells. The cells are used to produce an MTSP protein or protease domain thereof by growing the above-described cells under conditions whereby the encoded MTSP protein or protease domain of the MTSP protein is expressed by the cell, and recovering the expressed protease domain protein. For purposes herein, the protease domain is preferably secreted into the medium.

In one embodiment, the vectors include a sequence of nucleotides that encodes a polypeptide that has protease activity and contains all or a portion of only the protease domain, or multiple copies thereof, of an MTSP protein are provided. Also provided are vectors that comprise a sequence of nucleotides that encodes the protease domain and additional portions of an MTSP protein up to and including a full length MTSP protein, as well as multiple copies thereof, are also provided. The vectors may selected for expression of the MTSP protein or protease domain thereof in the cell or such that the MTSP protein is

WO 01/57194

5

10

-65-

PCT/US01/03471

expressed as a transmembrane protein. Alternatively, the vectors may include signals necessary for secretion of encoded proteins. When the protease domain is expressed the nucleic acid is preferably linked to a secretion signal, such as the *Saccharomyces cerevisiae*  $\alpha$  mating factor signal sequence or a portion thereof.

A variety of host-vector systems may be used to express the protein coding sequence. These include but are not limited to mammalian cell systems infected with virus (e.g. vaccinia virus, adenovirus, etc.); insect cell systems infected with virus (e.g. baculovirus); microorganisms such as yeast containing yeast vectors; or bacteria transformed with bacteriophage, DNA, plasmid DNA, or cosmid DNA. The expression elements of vectors vary in their strengths and specificities. Depending on the host-vector system used, any one of a number of suitable transcription and translation elements may be used.

Any methods known to those of skill in the art for the insertion of DNA 15 fragments into a vector may be used to construct expression vectors containing a chimeric gene containing appropriate transcriptional/translational control signals and protein coding sequences. These methods may include in vitro recombinant DNA and synthetic techniques and in vivo recombinants (genetic recombination). Expression of nucleic acid sequences encoding MTSP protein, or domains, 20 derivatives, fragments or homologs thereof, may be regulated by a second nucleic acid sequence so that the genes or fragments thereof are expressed in a host transformed with the recombinant DNA molecule(s). For example, expression of the proteins may be controlled by any promoter/enhancer known in the art. In a specific embodiment, the promoter is not native to the genes for MTSP protein. Promoters which may be used include but are not limited to the 25 SV40 early promoter (Bernoist and Chambon, Nature 290:304-310 (1981)), the promoter contained in the 3' long terminal repeat of Rous sarcoma virus (Yamamoto et al., Cell 22:787-797 (1980)), the herpes thymidine kinase promoter (Wagner et al., Proc. Natl. Acad. Sci. USA 78:1441-1445 (1981)), the regulatory sequences of the metallothionein gene (Brinster et al., Nature 296:39-30 42 (1982)); prokaryotic expression vectors such as the  $\beta$ -lactamase promoter (Villa-Kamaroff et al., Proc. Natl. Acad. Sci. USA 75:3727-3731 1978)) or the

tac promoter (DeBoer et al., Proc. Natl. Acad. Sci. USA 80:21-25 (1983)); see also "Useful Proteins from Recombinant Bacteria": in Scientific American 242:79-94 (1980)); plant expression vectors containing the nopaline synthetase promoter (Herrar-Estrella et al., Nature 303:209-213 (1984)) or the cauliflower mosaic virus 35S RNA promoter (Garder et al., Nucleic Acids Res. 9:2871 (1981)), and the promoter of the photosynthetic enzyme ribulose bisphosphate carboxylase (Herrera-Estrella et al., Nature 310:115-120 (1984)); promoter elements from yeast and other fungi such as the Gal4 promoter, the alcohol dehydrogenase promoter, the phosphoglycerol kinase promoter, the alkaline 10 phosphatase promoter, and the following animal transcriptional control regions that exhibit tissue specificity and have been used in transgenic animals: elastase I gene control region which is active in pancreatic acinar cells (Swift et al., Cell 38:639-646 (1984); Ornitz et al., Cold Spring Harbor Symp. Quant. Biol. 50:399-409 (1986); MacDonald, Hepatology 7:425-515 (1987)); insulin gene 15 control region which is active in pancreatic beta cells (Hanahan et al., Nature 315:115-122 (1985)), immunoglobulin gene control region which is active in lymphoid cells (Grosschedl et al., Cell 38:647-658 (1984); Adams et al., Nature 318:533-538 (1985); Alexander et al., Mol. Cell Biol. 7:1436-1444 (1987)), mouse mammary tumor virus control region which is active in testicular, breast, 20 lymphoid and mast cells (Leder et al., Cell 45:485-495 (1986)), albumin gene control region which is active in liver (Pinckert et al., Genes and Devel. 1:268-276 (1987)), alpha-fetoprotein gene control region which is active in liver (Krumlauf et al., Mol. Cell. Biol. 5:1639-1648 (1985); Hammer et al., Science 235:53-58 1987)), alpha-1 antitrypsin gene control region which is active in liver 25 (Kelsey et al., Genes and Devel. 1:161-171 (1987)), beta globin gene control region which is active in myeloid cells (Mogram et al., Nature 315:338-340 (1985); Kollias et al., Cell 46:89-94 (1986)), myelin basic protein gene control region which is active in oligodendrocyte cells of the brain (Readhead et al., Cell 48:703-712 (1987)), myosin light chain-2 gene control region which is active in 30 skeletal muscle (Sani, Nature 314:283-286 (1985)), and gonadotrophic releasing hormone gene control region which is active in gonadotrophs of the hypothalamus (Mason et al., Science 234:1372-1378 (1986)).

-67-

In a specific embodiment, a vector is used that contains a promoter operably linked to nucleic acids encoding an MTSP protein, or a domain, fragment, derivative or homolog, thereof, one or more origins of replication, and optionally, one or more selectable markers (*e.g.*, an antibiotic resistance gene). Expression vectors containing the coding sequences, or portions thereof, of an MTSP protein, is made, for example, by subcloning the coding portions into the EcoRI restriction site of each of the three pGEX vectors (glutathione Stransferase expression vectors (Smith and Johnson, *Gene* 7:31-40 (1988)). This allows for the expression of products in the correct reading frame. Preferred vectors and systems for expression of the protease domains of the MTSP proteins are well known *Pichia* vectors (available, for example, from Invitrogen, San Diego, CA), particularly those designed for secretion of the encoded proteins. One exemplary vector is described in the EXAMPLES.

10

15

20

25

30

Plasmids for transformation of <u>E. coli</u> cells, include, for example, the pET expression vectors (see, U.S patent 4,952,496; available from NOVAGEN, Madison, WI; see, also literature published by Novagen describing the system). Such plasmids include pET 11a, which contains the T7lac promoter, T7 terminator, the inducible <u>E. coli</u> lac operator, and the lac repressor gene; pET 12a-c, which contains the T7 promoter, T7 terminator, and the <u>E. coli</u> ompT secretion signal; and pET 15b and pET19b (NOVAGEN, Madison, WI), which contain a His-Tag<sup>™</sup> leader sequence for use in purification with a His column and a thrombin cleavage site that permits cleavage following purification over the column; the T7-lac promoter region and the T7 terminator.

The vectors are introduced into host cells, such as *Pichia* cells and bacterial cells, such as *E. coli*, and the proteins expressed therein. Preferred *Pichia* strains, include, for example, GS115. Preferred bacterial hosts contain chromosomal copies of DNA encoding T7 RNA polymerase operably linked to an inducible promoter, such as the lacUV promoter (see, U.S. Patent No. 4,952,496). Such hosts include, but are not limited to, the lysogenic <u>E. coli</u> strain BL21(DE3).

-68-

# Expression and production of proteins

5

10

15

20

25

30

The MTSP domains, derivatives and analogs be produced by various methods known in the art. For example, once a recombinant cell expressing an MTSP protein, or a domain, fragment or derivative thereof, is identified, the individual gene product can be isolated and analyzed. This is achieved by assays based on the physical and/or functional properties of the protein, including, but not limited to, radioactive labeling of the product followed by analysis by gel electrophoresis, immunoassay, cross-linking to marker-labeled product. The MTSP protein proteins may be isolated and purified by standard methods known in the art (either from natural sources or recombinant host cells expressing the complexes or proteins), including but not restricted to column chromatography (e.g., ion exchange, affinity, gel exclusion, reversed-phase high pressure, fast protein liquid, etc.), differential centrifugation, differential solubility, or by any other standard technique used for the purification of proteins. Functional properties may be evaluated using any suitable assay known in the art.

Alternatively, once an MTSP protein or its domain or derivative is identified, the amino acid sequence of the protein can be deduced from the nucleotide sequence of the gene which encodes it. As a result, the protein or its domain or derivative can be synthesized by standard chemical methods known in the art (e.g. see Hunkapiller et al, Nature 310:105-111 (1984)).

Manipulations of MTSP protein sequences may be made at the protein level. Also contemplated herein are MTSP protein proteins, domains thereof, derivatives or analogs or fragments thereof, which are differentially modified during or after translation, *e.g.*, by glycosylation, acetylation, phosphorylation, amidation, derivatization by known protecting/blocking groups, proteolytic cleavage, linkage to an antibody molecule or other cellular ligand, etc. Any of numerous chemical modifications may be carried out by known techniques, including but not limited to specific chemical cleavage by cyanogen bromide, trypsin, chymotrypsin, papain, V8 protease, NaBH<sub>4</sub>, acetylation, formylation, oxidation, reduction, metabolic synthesis in the presence of tunicamycin, etc.

-69-

In addition, domains, analogs and derivatives of an MTSP protein can be chemically synthesized. For example, a peptide corresponding to a portion of an MTSP protein, which includes the desired domain or which mediates the desired activity *in vitro* can be synthesized by use of a peptide synthesizer.

Furthermore, if desired, nonclassical amino acids or chemical amino acid analogs can be introduced as a substitution or addition into the MTSP protein sequence. Non-classical amino acids include but are not limited to the D-isomers of the common amino acids, a-amino isobutyric acid, 4-aminobutyric acid, Abu, 2-aminobutyric acid, e-Abu, e-Ahx, 6-amino hexanoic acid, Aib, 2-amino isobutyric acid, 3-amino propionoic acid, ornithine, norleucine, norvaline, hydroxyproline, sarcosine, citrulline, cysteic acid, t-butylglycine, t-butylalanine, phenylglycine, cyclohexylalanine, ß-alanine, fluoro-amino acids, designer amino acids such as ß-methyl amino acids, Ca-methyl amino acids, Na-methyl amino acids, and amino acid analogs in general. Furthermore, the amino acid can be D (dextrorotary) or L (levorotary).

In cases where natural products are suspected of being mutant or are isolated from new species, the amino acid sequence of the MTSP protein isolated from the natural source, as well as those expressed *in vitro*, or from synthesized expression vectors *in vivo* or *in vitro*, can be determined from analysis of the DNA sequence, or alternatively, by direct sequencing of the isolated protein. Such analysis may be performed by manual sequencing or through use of an automated amino acid sequenator.

#### Modifications

10

15

20

25

30

A variety of modification of the MTSP proteins and domains are contemplated herein. An MTSP-encoding nucleic acid molecule modified by any of numerous strategies known in the art (Sambrook et al., 1990, *Molecular Cloning, A Laboratory Manual*, 2d ed., Cold Spring Harbor Laboratory, Cold Spring Harbor, New York). The sequences can be cleaved at appropriate sites with restriction endonuclease(s), followed by further enzymatic modification if desired, isolated, and ligated *in vitro*. In the production of the gene encoding a domain, derivative or analog of MTSP, care should be taken to ensure that the modified gene retains the original translational reading frame, uninterrupted by

translational stop signals, in the gene region where the desired activity is encoded.

Additionally, the MTSP-encoding nucleic acid molecules can be mutated in vitro or in vivo, to create and/or destroy translation, initiation, and/or termination sequences, or to create variations in coding regions and/or form new restriction endonuclease sites or destroy pre-existing ones, to facilitate further in vitro modification. Also, as described herein muteins with primary sequence alterations, such as replacements of Cys residues and elimination of glycosylation sites are contemplated. Such mutations may be effected by any technique for mutagenesis known in the art, including, but not limited to, chemical mutagenesis and in vitro site-directed mutagenesis (Hutchinson et al., J. Biol. Chem. 253:6551-6558 (1978)), use of TAB® linkers (Pharmacia). In one embodiment, for example, an MTSP protein or domain thereof is modified to include a fluorescent label. In other specific embodiments, the MTSP protein is modified to have a heterofunctional reagent, such heterofunctional reagents can be used to crosslink the members of the complex.

10

15

20

25

30

The MTSP proteins may be isolated and purified by standard methods known in the art (either from natural sources or recombinant host cells expressing the complexes or proteins), including but not restricted to column chromatography (e.g., ion exchange, affinity, gel exclusion, reversed-phase high pressure, fast protein liquid, etc.), differential centrifugation, differential solubility, or by any other standard technique used for the purification of proteins. Functional properties may be evaluated using any suitable assay known in the art.

Alternatively, once a MTSP or its domain or derivative is identified, the amino acid sequence of the protein can be deduced from the nucleotide sequence of the gene which encodes it. As a result, the protein or its domain or derivative can be synthesized by standard chemical methods known in the art (e.g., see Hunkapiller et al, *Nature*, 310:105-111 (1984)).

Manipulations of MTSP sequences may be made at the protein level. MTSP domains, derivatives or analogs or fragments, which are differentially modified during or after translation, *e.g.*, by glycosylation, acetylation,

-71-

phosphorylation, amidation, derivatization by known protecting/blocking groups, proteolytic cleavage, linkage to an antibody molecule and other cellular ligand, are contemplated herein. Any of numerous chemical modifications may be carried out by known techniques, including but not limited to specific chemical cleavage by cyanogen bromide, trypsin, chymotrypsin, papain, V8 protease, NaBH<sub>4</sub>, acetylation, formylation, oxidation, reduction, metabolic synthesis in the presence of tunicamycin, etc.

In addition, domains, analogs and derivatives of a MTSP can be chemically synthesized. For example, a peptide corresponding to a portion of a MTSP, which comprises the desired domain or which mediates the desired activity *in vitro* can be synthesized by use of a peptide synthesizer. Furthermore, if desired, nonclassical amino acids or chemical amino acid analogs can be introduced as a substitution or addition into the MTSP sequence. Nonclassical amino acids include but are not limited to the D-isomers of the common amino acids, a-amino isobutyric acid, 4-aminobutyric acid, Abu, 2-aminobutyric acid, 3-amino propionoic acid, ornithine, norleucine, norvaline, hydroxyproline, sarcosine, citrulline, cysteic acid, t-butylglycine, t-butylalanine, phenylglycine, cyclohexylalanine,  $\mathcal{B}$ -alanine, fluoro-amino acids, designer amino acids such as  $\mathcal{B}$ -methyl amino acids, Ca-methyl amino acids, Na-methyl amino acids, and amino acid analogs in general. Furthermore, the amino acid can be D (dextrorotary) or L (levorotary).

#### F. SCREENING METHODS

5

10

15

20

25

30

The single chain protease domains, as shown herein, can be used in a variety of methods to identify compounds that modulate the activity thereof. For MTSPs that exhibit higher activity or expression in tumor cells, compounds that inhibit the proteolytic activity are of particular interest. For any MTSPs that are active at lower levels in tumor cells, compounds or agents that enhance the activity are potentially of interest. In all instances the identified compounds will include agents that are candidate cancer treatments.

Several types of assays are exemplified and described herein. It is understood that the protease domains may be used in other assays. It is shown

-72-

here, however, that the single chain protease domains exhibit catalytic activity. As such they are ideal for *in vitro* screening assays.

They may also be used in binding assays.

10

15

20

25

30

The MTSP3, MTSP4 and MTSP6 full length zymogens, activated enzymes, single and double chain protease domains are contemplated for use in any screening assay known to those of skill in the art, including those provided herein. Hence the following description, if directed to proteolytic assays is intended to apply to use of a single chain protease domain or a catalytically active portion thereof of any MTSP, including an MTSP3, MTSP4 or an MTSP6. Other assays, such as binding assays are provided herein, particularly for use with an MTSP3, MTSP4 or MTSP6, including any variants, such as splice variants thereof. MTSP3 and MTSP4 are of most interest in such assays.

# 1. Catalytic Assays for identification of agents that modulate the protease activity of an MTSP protein

Methods for identifying a modulator of the catalytic activity of an MTSP, particularly a single chain protease domain or catalytically active portion thereof, are provided herein. The methods can be practiced by: a) contacting the MTSP, particularly a single-chain domain thereof, with a substrate of the MTSP in the presence of a test substance, and detecting the proteolysis of the substrate, whereby the activity of the MTSP is assessed, and comparing the activity to a control. For example, the control can be the activity of the MTSP assessed by contacting an MTSP, particularly a single-chain domain thereof, with a substrate of the MTSP, and detecting the proteolysis of the substrate, whereby the activity of the MTSP is assessed. The results in the presence and absence of the test compounds are compared. A difference in the activity indicates that the test substance modulates the activity of the MTSP.

In one embodiment a plurality of the test substances are screened simultaneously in the above screening method. In another embodiment, the MTSP is isolated from a target cell as a means for then identifying agents that are potentially specific for the target cell.

In still another embodiment, The test substance is a therapeutic compound, and whereby a difference of the MTSP activity measured in the

presence and in the absence of the test substance indicates that the target cell responds to the therapeutic compound.

One method include the steps of (a) contacting the MTSP protein or protease domain thereof with one or a plurality of test compounds under conditions conducive to interaction between the ligand and the compounds; and (b) identifying one or more compounds in the plurality that specifically binds to the ligand.

5

10

15

20

25

30

Another method provided herein includes the steps of a) contacting an MTSP protein or protease domain thereof with a substrate of the MTSP protein, and detecting the proteolysis of the substrate, whereby the activity of the MTSP protein is assessed; b) contacting the MTSP protein with a substrate of the MTSP protein in the presence of a test substance, and detecting the proteolysis of the substrate, whereby the activity of the MTSP protein is assessed; and c) comparing the activity of the MTSP protein assessed in steps a) and b), whereby the activity measured in step a) differs from the activity measured in step b) indicates that the test substance modulates the activity of the MTSP protein.

In another embodiment, a plurality of the test substances are screened simultaneously. In comparing the activity of an MTSP protein in the presence and absence of a test substance to assess whether the test substance is a modulator of the MTSP protein, it is unnecessary to assay the activity in parallel, although such parallel measurement is preferred. It is possible to measure the activity of the MTSP protein at one time point and compare the measured activity to a historical value of the activity of the MTSP protein.

For instance, one can measure the activity of the MTSP protein in the presence of a test substance and compare with historical value of the activity of the MTSP protein measured previously in the absence of the test substance, and *vice versa*. This can be accomplished, for example, by providing the activity of the MTSP protein on an insert or pamphlet provided with a kit for conducting the assay.

Methods for selecting substrates for a particular MTSP are described in the EXAMPLES, and particular proteolytic assays are exemplified.

-74-

Combinations and kits containing the combinations optionally including instructions for performing the assays are provided. The combinations include an MTSP protein and a substrate of the MTSP protein to be assayed; and, optionally reagents for detecting proteolysis of the substrate. The substrates, which are typically proteins subject to proteolysis by a particular MTSP protein, can be identified empirically by testing the ability of the MTSP protein to cleave the test substrate. Substrates that are cleaved most effectively (i.e., at the lowest concentrations and/or fastest rate or under desirable conditions), are identified.

Additionally provided herein is a kit containing the above-described combination. Preferably, the kit further includes instructions for identifying a modulator of the activity of an MTSP protein. Any MTSP protein is contemplated as target for identifying modulators of the activity thereof.

#### 2. Binding assays

10

15

20

25

30

Also provided herein are methods for identification and isolation of agents, particularly compounds that bind to MTSPs. The assays are designed to identify agents that bind to the zymogen form, the single chain isolated protease domain (or a protein, other than an MTSP protein, that contains the protease domain of an MTSP protein), and to the activated form, including the activated form derived from the full length zymogen or from an extended protease domain. The identified compounds are candidates or leads for identification of compounds for treatments of tumors and other disorders and diseases involving aberrant angiogenesis. The MTSP proteins used in the methods include any MTSP protein as defined herein, and preferably use MTSP single chain domain or proteolytically active portion thereof.

A variety of methods are provided herein. These methods may be performed in solution or in solid phase reactions in which the MTSP protein(s) or protease domain(s) thereof are linked, either directly or indirectly via a linker, to a solid support. Screening assays are described in the Examples, and these assays have been used to identify candidate compounds.

For purposes herein, all binding assays described above are provided for MTSP3, MTSP4 and MTSP6. For MTSP1 (including any variant thereof) and

thereof) and other such proteases, binding assays that employ the isolated single chain protease domain or a protein containing such domain (other than the MTSP from which the protease is derived) are provided.

Methods for identifying an agent, such as a compound, that specifically binds to an MTSP single chain protease domain or an MTSP, such as an MTSP3, MTSP4 or an MTSP6, are provided herein. The method can be practiced by (a) contacting the MTSP with one or a plurality of test agents under conditions conducive to binding between the MTSP and an agent; and (b) identifying one or more agents within the plurality that specifically binds to the MTSP.

5

10

15

20

25

For example, in practicing such methods the MTSP polypeptide is mixed with a potential binding partner or an extract or fraction of a cell under conditions that allow the association of potential binding partners with the polypeptide. After mixing, peptides, polypeptides, proteins or other molecules that have become associated with an MTSP are separated from the mixture. The binding partner that bound to the MTSP can then be removed and further analyzed. To identify and isolate a binding partner, the entire protein, for instance the entire disclosed protein of SEQ ID Nos. 6, 8 10 or 12 can be used. Alternatively, a fragment of the protein can be used.

A variety of methods can be used to obtain cell extracts. Cells can be disrupted using either physical or chemical disruption methods. Examples of physical disruption methods include, but are not limited to, sonication and mechanical shearing. Examples of chemical lysis methods include, but are not limited to, detergent lysis and enzyme lysis. A skilled artisan can readily adapt methods for preparing cellular extracts in order to obtain extracts for use in the present methods.

Once an extract of a cell is prepared, the extract is mixed with the MTSP under conditions in which association of the protein with the binding partner can occur. A variety of conditions can be used, the most preferred being conditions that closely resemble conditions found in the cytoplasm of a human cell.

30 Features such as osmolarity, pH, temperature, and the concentration of cellular extract used, can be varied to optimize the association of the protein with the binding partner.

After mixing under appropriate conditions, the bound complex is separated from the mixture. A variety of techniques can be used to separate the mixture. For example, antibodies specific to an MTSP can be used to immunoprecipitate the binding partner complex. Alternatively, standard chemical separation techniques such as chromatography and density/sediment centrifugation can be used.

5

10

15

20

25

30

After removing the non-associated cellular constituents in the extract, the binding partner can be dissociated from the complex using conventional methods. For example, dissociation can be accomplished by altering the salt concentration or pH of the mixture.

To aid in separating associated binding partner pairs from the mixed extract, the MTSP can be immobilized on a solid support. For example, the protein can be attached to a nitrocellulose matrix or acrylic beads. Attachment of the protein or a fragment thereof to a solid support aids in separating peptide/binding partner pairs from other constituents found in the extract. The identified binding partners can be either a single protein or a complex made up of two or more proteins.

Alternatively, the nucleic acid molecules encoding the single chain proteases can be used in a yeast two-hybrid system. The yeast two-hybrid system has been used to identify other protein partner pairs and can readily be adapted to employ the nucleic acid molecules herein described.

Another *in vitro* binding assay, particularly for an MTSP3, MTSP4 or an MTSP6 uses a mixture of a polypeptide that contains at least the catalytic domain of one of these proteins and one or more candidate binding targets or substrates. After incubating the mixture under appropriate conditions, one determines whether the MTSP or a polypeptide fragment thereof containing the catalytic domain binds with the candidate substrate. For cell-free binding assays, one of the components includes or is coupled to a detectable label. The label may provide for direct detection, such as radioactivity, luminescence, optical or electron density, *etc.*, or indirect detection such as an epitope tag, an enzyme, *etc.* A variety of methods may be employed to detect the label depending on the nature of the label and other assay components. For example,

*-*77-

the label may be detected bound to the solid substrate or a portion of the bound complex containing the label may be separated from the solid substrate, and the label thereafter detected.

#### 3. Detection of signal transduction

5

10

15

20

25

30

The cell surface location of the MTSPs suggests a role for some or all of these proteins in signal transduction. Assays for assessing signal transduction are well known to those of skill in the art, and may be adapted for use with the MTSP protein.

Assays for identifying agents that effect or alter signal transduction mediated by an MTSP, particularly the full length or a sufficient portion to anchor the extracellular domain or a function portion thereof of an MTSP on the surface of a cell are provided. Such assays, include, for example, transcription based assays in which modulation of a transduced signal is assessed by detecting an effect on an expression from a reporter gene (see, *e.g.*, U.S. Patent No. 5,436,128).

# 4. Methods for Identifying Agents that Modulate the Expression a Nucleic Acid Encoding an MTSP, particularly an MTSP3, MTSP4 or MTSP6

Another embodiment provides methods for identifying agents that modulate the expression of a nucleic acid encoding an MTSP, particularly an MTSP3, MTSP4 or MTSP. Such assays use any available means of monitoring for changes in the expression level of the nucleic acids encoding an MTSP, such as MTSP3 or MTSP4.

In one assay format, cell lines that contain reporter gene fusions between the open reading frame of MTSP3, MTSP4 or MTSP6 or a domain thereof, particularly the protease domain and any assayable fusion partner may be prepared. Numerous assayable fusion partners are known and readily available including the firefly luciferase gene and the gene encoding chloramphenicol acetyltransferase (Alam *et al.*, *Anal. Biochem*. 188: 245-54 (1990)). Cell lines containing the reporter gene fusions are then exposed to the agent to be tested under appropriate conditions and time. Differential expression of the reporter gene between samples exposed to the agent and control samples identifies

-78-

agents which modulate the expression of a nucleic acid encoding an MTSP3, MTSP4 or MTSP6.

Additional assay formats may be used to monitor the ability of the agent to modulate the expression of a nucleic acid encoding an MTSP3, MTSP4 or MTSP6. For instance, mRNA expression may be monitored directly by hybridization to the nucleic acids. Cell lines are exposed to the agent to be tested under appropriate conditions and time and total RNA or mRNA is isolated by standard procedures (see, e.g., Sambrook et al. (1989) MOLECULAR CLONING: A LABORATORY MANUAL, 2nd Ed. Cold Spring Harbor Laboratory Press). Probes to detect differences in RNA expression levels between cells 10 exposed to the agent and control cells may be prepared from the nucleic acids. It is preferable, but not necessary, to design probes which hybridize only with target nucleic acids under conditions of high stringency. Only highly complementary nucleic acid hybrids form under conditions of high stringency. 15 Accordingly, the stringency of the assay conditions determines the amount of complementarity which should exist between two nucleic acid strands in order to form a hybrid. Stringency should be chosen to maximize the difference in stability between the probe:target hybrid and potential probe:non-target hybrids.

Probes may be designed from the nucleic acids through methods known in the art. For instance, the G+C content of the probe and the probe length can affect probe binding to its target sequence. Methods to optimize probe specificity are commonly available (see, e.g., Sambrook et al. (1989) MOLECULAR CLONING: A LABORATORY MANUAL, 2nd Ed. Cold Spring Harbor Laboratory Press); and Ausubel et al. (1995) CURRENT PROTOCOLS IN MOLECULAR BIOLOGY, Greene Publishing Co., NY).

20

25

30

Hybridization conditions are modified using known methods (see, *e.g.*, Sambrook *et al.* (1989) MOLECULAR CLONING: A LABORATORY MANUAL, 2nd Ed. Cold Spring Harbor Laboratory Press); and Ausubel *et al.* (1995) CURRENT PROTOCOLS IN MOLECULAR BIOLOGY, Greene Publishing Co., NY), as required for each probe. Hybridization of total cellular RNA or RNA enriched for polyA RNA can be accomplished in any available format. For instance, total cellular RNA or RNA enriched for polyA RNA can be affixed to a solid support,

and the solid support exposed to at least one probe comprising at least one, or part of one of the nucleic acid molecules under conditions in which the probe will specifically hybridize. Alternatively, nucleic acid fragments comprising at least one, or part of one of the sequences can be affixed to a solid support, such as a porous glass wafer. The glass wafer can then be exposed to total cellular RNA or polyA RNA from a sample under conditions in which the affixed sequences will specifically hybridize. Such glass wafers and hybridization methods are widely available, for example, those disclosed by Beattie (WO 95/11755). By examining for the ability of a given probe to specifically hybridize to an RNA sample from an untreated cell population and from a cell population exposed to the agent, agents which up or down regulate the expression of a nucleic acid encoding the protein having the sequence of SEQ ID NO:3 or SEQ ID NO:4 are identified.

5

10

15

20

25

30

5. Methods for Identifying Agents that Modulate at Least One Activity of an MTPS, such as MTSP3, MTSP4 or MTSP6

Methods for identifying agents that modulate at least one activity of a an MTSP, such as an MTSP3, MTSP4 or MTSP6 are provided. Such methods or assays may use any means of monitoring or detecting the desired activity.

In one format, the relative amounts of a protein between a cell population that has been exposed to the agent to be tested compared to an un-exposed control cell population may be assayed (e.g., a prostate cancer cell line, a lung cancer cell line, a colon cancer cell line or a breast cancer cell line). In this format, probes, such as specific antibodies, are used to monitor the differential expression of the protein in the different cell populations. Cell lines or populations are exposed to the agent to be tested under appropriate conditions and time. Cellular lysates may be prepared from the exposed cell line or population and a control, unexposed cell line or population. The cellular lysates are then analyzed with the probe.

For example, N- and C- terminal fragments of the MTSP can be expressed in bacteria and used to search for proteins which bind to these fragments.

Fusion proteins, such as His-tag or GST fusion to the N- or C-terminal regions of the MTSP, such as an MTSP3, MTSP4 or an MTSP6, can be prepared for use as

-80-

a substrate. These fusion proteins can be coupled to, for example, Glutathione-Sepharose beads and then probed with cell lysates. Prior to lysis, the cells may be treated with a candidate agent which may modulate an MTSP, such as an MTSP3, MTSP4 or an MTSP6, or proteins that interact with domains thereon.

Lysate proteins binding to the fusion proteins can be resolved by SDS-PAGE, isolated and identified by protein sequencing or mass spectroscopy, as is known in the art.

10

15

20

25

30

Antibody probes are prepared by immunizing suitable mammalian hosts in appropriate immunization protocols using the peptides, polypeptides or proteins if they are of sufficient length (e.g., 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 20, 25, 30, 35, 40 or more consecutive amino acids the MTSP protein, such as an MTSP3, an MTSP4 or an MTSP6), or if required to enhance immunogenicity, conjugated to suitable carriers. Methods for preparing immunogenic conjugates with carriers, such as bovine serum albumin (BSA), keyhole limpet hemocyanin (KLH), or other carrier proteins are well known in the art. In some circumstances, direct conjugation using, for example, carbodiimide reagents may be effective; in other instances linking reagents such as those supplied by Pierce Chemical Co., Rockford, IL, may be desirable to provide accessibility to the hapten. Hapten peptides can be extended at either the amino or carboxy terminus with a Cys residue or interspersed with cysteine residues, for example, to facilitate linking to a carrier. Administration of the immunogens is conducted generally by injection over a suitable time period and with use of suitable adjuvants, as is generally understood in the art. During the immunization schedule, titers of antibodies are taken to determine adequacy of antibody formation.

Anti-peptide antibodies can be generated using synthetic peptides corresponding to, for example, the carboxy terminal amino acids of the MTSP. Synthetic peptides can be as small as 1-3 amino acids in length, but are preferably at least 4 or more amino acid residues long. The peptides can be coupled to KLH using standard methods and can be immunized into animals, such as rabbits or ungulate. Polyclonal antibodies can then be purified, for example using Actigel beads containing the covalently bound peptide.

While the polyclonal antisera produced in this way may be satisfactory for some applications, for pharmaceutical compositions, use of monoclonal preparations is preferred. Immortalized cell lines which secrete the desired monoclonal antibodies may be prepared using the standard method of Kohler *et al.*, (*Nature* 256: 495-7 (1975)) or modifications which effect immortalization of lymphocytes or spleen cells, as is generally known. The immortalized cell lines secreting the desired antibodies are screened by immunoassay in which the antigen is the peptide hapten, polypeptide or protein. When the appropriate immortalized cell culture secreting the desired antibody is identified, the cells can be cultured either *in vitro* or by production *in vivo* via ascites fluid. Of particular interest, are monoclonal antibodies that recognize the catalytic domain of an MTSP, such as an MTSP3, MTSP4 or an MTSP6.

10

15

20

25

30

Additionally, the zymogen or two-chain forms the MTSP can be used to make monoclonal antibodies which recognize conformation epitopes. For peptide-directed monoclonal antibodies, peptides from the C1r/C1s domain can be used to generate anti-C1r/C1s domain monoclonal antibodies which can thereby block activation of the zymogen to the two-chain form of the MTSP. This domain can similarly be the substrate for other non-antibody compounds which bind to these preferred domains on either the single-chain or double-chain forms of the MTSP3, MTSP4 or MTSP6, and thereby modulate the activity of thereof or prevent its activation.

The desired monoclonal antibodies are then recovered from the culture supernatant or from the ascites supernatant. Fragments of the monoclonals or the polyclonal antisera which contain the immunologically significant portion can be used as antagonists, as well as the intact antibodies. Use of immunologically reactive fragments, such as the Fab, Fab', of F(ab')<sub>2</sub> fragments are often preferable, especially in a therapeutic context, as these fragments are generally less immunogenic than the whole immunoglobulin.

The antibodies or fragments may also be produced. Regions that bind specifically to the desired regions of receptor can also be produced in the context of chimeras with multiple species origin.

-82-

Agents that are assayed in the above method can be randomly selected or rationally selected or designed.

The agents can be, as examples, peptides, small molecules, and carbohydrates. A skilled artisan can readily recognize that there is no limit as to the structural nature of the agents.

The peptide agents can be prepared using standard solid phase (or solution phase) peptide synthesis methods, as is known in the art. In addition, the DNA encoding these peptides may be synthesized using commercially available oligonucleotide synthesis instrumentation and produced recombinantly using standard recombinant production systems. The production using solid phase peptide synthesis is necessitated if non-gene-encoded amino acids are to be included.

#### G. Assay formats and selection of test substances

5

10

15

A variety of formats and detection protocols are known for performing screening assays. Any such formats and protocols may be adapted for identifying modulators of MTSP protein activities. The following includes a discussion of exemplary protocols.

#### 1. High throughput screening assays

Although the above-described assay can be conducted where a single

MTSP protein is screened, and/or a single test substance is screened for in one assay, the assay is preferably conducted in a high throughput screening mode, i.e., a plurality of the MTSP proteins are screened against and/or a plurality of the test substances are screened for simultaneously (See generally, High Throughput Screening: The Discovery of Bioactive Substances (Devlin, Ed.)

Marcel Dekker, 1997; Sittampalam et al., Curr. Opin. Chem. Biol., 1(3):384-91 (1997); and Silverman et al., Curr. Opin. Chem. Biol., 2(3):397-403 (1998)). For example, the assay can be conducted in a multi-well (e.g., 24-, 48-, 96-, or 384-well), chip or array format.

High-throughput screening (HTS) is the process of testing a large number of diverse chemical structures against disease targets to identify "hits" (Sittampalam et al., *Curr. Opin. Chem. Biol.*, 1(3):384-91 (1997)). Current state-of-the-art HTS operations are highly automated and computerized to handle

sample preparation, assay procedures and the subsequent processing of large volumes of data.

Detection technologies employed in high-throughput screens depend on the type of biochemical pathway being investigated (Sittampalam et al., Curr. Opin. Chem. Biol., 1(3):384-91 (1997)). These methods include, radiochemical 5 methods, such as the scintillation proximity assays (SPA), which can be adapted to a variety of enzyme assays (Lerner et al., J. Biomol. Screening, 1:135-143 (1996); Baker et al., Anal. Biochem., 239:20-24 (1996); Baum et al., Anal. Biochem., 237:129-134 (1996); and Sullivan et al., J. Biomol. Screening, 2:19-23 (1997)) and protein-protein interaction assays (Braunwalder et al., J. Biomol. 10 Screening, 1:23-26 (1996); Sonatore et al., Anal. Biochem., 240:289-297 (1996); and Chen et al., J. Biol. Chem., 271:25308-25315 (1996)), and nonisotopic detection methods, including but are not limited to, colorimetric and luminescence detection methods, resonance energy transfer (RET) methods, 15 time-resolved fluorescence (HTRF) methods, cell-based fluorescence assays, such as fluorescence resonance energy transfer (FRET) procedures (see, e.g., Gonzalez et al., Biophys. J., 69:1272-1280 (1995)), fluorescence polarization or anisotropy methods (see, e.g., Jameson et al., Methods Enzymol., 246:283-300 (1995); Jolley, J. Biomol. Screening, 1:33-38 (1996); Lynch et al., Anal. Biochem., 247:77-82 (1997)), fluorescence correlation spectroscopy (FCS) 20 and other such methods.

#### 2. Test Substances

25

30

Test compounds, including small molecules and libraries and collections thereof can be screened in the above-described assays and assays described below to identify compounds that modulate the activity an MTSP protein. Rational drug design methodologies that rely on computational chemistry may be used to screen and identify candidate compounds.

The compounds identified by the screening methods include inhibitors, including antagonists, and may be agonists. Compounds for screening are any compounds and collections of compounds available, know or that can be prepared.

-84-

#### a. Selection of Compounds

5

10

15

20

25

30

Compounds can be selected for their potency and selectivity of inhibition of serine proteases, especially MTSP protein. As described herein, and as generally known, a target serine protease and its substrate are combined under assay conditions permitting reaction of the protease with its substrate. The assay is performed in the absence of test compound, and in the presence of increasing concentrations of the test compound. The concentration of test compound at which 50% of the serine protease activity is inhibited by the test compound is the IC $_{50}$  value (Inhibitory Concentration) or EC $_{50}$  (Effective Concentration) value for that compound. Within a series or group of test compounds, those having lower IC $_{50}$  or EC $_{50}$  values are considered more potent inhibitors of the serine protease than those compounds having higher IC $_{50}$  or EC $_{50}$  values. The IC $_{50}$  measurement is often used for more simplistic assays, whereas the EC $_{50}$  is often used for more complicated assays, such as those employing cells.

Preferred compounds according to this aspect have an  $IC_{50}$  value of 100 nM or less as measured in an *in vitro* assay for inhibition of MTSP protein activity. Especially preferred compounds have an  $IC_{50}$  value of less than 100 nM.

The test compounds also are evaluated for selectivity toward a serine protease. As described herein, and as generally known, a test compound is assayed for its potency toward a panel of serine proteases and other enzymes and an  $IC_{50}$  value or  $EC_{50}$  value is determined for each test compound in each assay system. A compound that demonstrates a low  $IC_{50}$  value or  $EC_{50}$  value for the target enzyme, *e.g.*, MTSP protein, and a higher  $IC_{50}$  value or  $EC_{50}$  value for other enzymes within the test panel (*e.g.*, urokinase tissue plasminogen activator, thrombin, Factor Xa), is considered to be selective toward the target enzyme. Generally, a compound is deemed selective if its  $IC_{50}$  value or  $EC_{50}$  value in the target enzyme assay is at least one order of magnitude less than the next smallest  $IC_{50}$  value or  $EC_{50}$  value measured in the selectivity panel of enzymes.

-85-

Presently preferred compounds have an  $IC_{50}$  value of 100 nM or less as measured in an *in vitro* assay for inhibition of urokinase activity. Especially preferred compounds have an  $IC_{50}$  value in the *in vitro* urokinase inhibition assay that is at least one order of magnitude smaller than the  $IC_{50}$  value measured in the *in vitro* tPA inhibition assay. Compounds having a selectivity ratio of  $IC_{50}$  u-PA assay:  $IC_{50}$  MTSP protein assay of greater than 100 are especially preferred.

Compounds are also evaluated for their activity *in vivo*. The type of assay chosen for evaluation of test compounds will depend on the pathological condition to be treated or prevented by use of the compound, as well as the route of administration to be evaluated for the test compound.

10

15

20

25

30

For instance, to evaluate the activity of a compound to reduce tumor growth through inhibition of MTSP protein, the procedures described by Jankun et al., *Canc. Res.*, <u>57</u>:559-563 (1997) to evaluate PAI-1 can be employed. Briefly, the ATCC cell lines DU145 and LnCaP are injected into SCID mice. After tumors are established, the mice are given test compound according to a dosing regime determined from the compound's *in vitro* characteristics. The Jankun *et al.* compound was administered in water. Tumor volume measurements are taken twice a week for about five weeks. A compound is deemed active if an animal to which the compound was administered exhibited decreased tumor volume, as compared to animals receiving appropriate control compounds.

Another *in vivo* experimental model designed to evaluate the effect of paraminobenzamidine, a swine protease inhibitor, on reducing tumor volume is described by Billström et al., *Int. J. Cancer*, <u>61</u>:542-547 (1995).

To evaluate the ability of a compound to reduce the occurrence of, or inhibit, metastasis, the procedures described by Kobayashi et al., *Int. J. Canc.*, <u>57</u>:727-733d (1994) can be employed. Briefly, a murein xenograft selected for high lung colonization potential in injected into C57B1/6 mice i.v. (experimental metastasis) or s.c. into the abdominal wall (spontaneous metastasis). Various concentrations of the compound to be tested can be admixed with the tumor cells in Matrigel prior to injection. Daily i.p. injections of the test compound are made either on days 1-6 or days 7-13 after tumor inoculation. The animals are sacrificed about three or four weeks after tumor inoculation, and the lung tumor

-86-

colonies are counted. Evaluation of the resulting data permits a determination as to efficacy of the test compound, optimal dosing and route of administration.

The activity of the tested compounds toward decreasing tumor volume and metastasis can be evaluated in model described in Rabbani et al., *Int. J. Cancer* 63:840-845 (1995) to evaluate their inhibitor. There, Mat LyLu tumor cells were injected into the flank of Copenhagen rats. The animals were implanted with osmotic minipumps to continuously administer various doses of test compound for up to three weeks. The tumor mass and volume of experimental and control animals were evaluated during the experiment, as were metastatic growths. Evaluation of the resulting data permits a determination as to efficacy of the test compound, optimal dosing, and route of administration. Some of these authors described a related protocol in Xing et al., *Canc. Res.*, 57:3585-3593 (1997).

5

10

15

20

25

30

To evaluate the inhibitory activity of a compound, a rabbit cornea neovascularization model can be employed. Avery et al., *Arch. Ophthalmol.*, 108:1474-1475 (1990) describe anesthetizing New Zealand albino rabbits and then making a central corneal incision and forming a radial corneal pocket. A slow release prostaglandin pellet was placed in the pocket to induce neovascularization. Test compound was administered i.p. for five days, at which time the animals were sacrificed. The effect of the test compound is evaluated by review of periodic photographs taken of the limbus, which can be used to calculate the area of neovascular response and, therefore, limbal neovascularization. A decreased area of neovascularization as compared with appropriate controls indicates the test compound was effective at decreasing or inhibiting neovascularization.

An angiogenesis model used to evaluate the effect of a test compound in preventing angiogenesis is described by Min et al., Canc. Res., 56:2428-2433 (1996). C57BL6 mice receive subcutaneous injections of a Matrigel mixture containing bFGF, as the angiogenesis-inducing agent, with and without the test compound. After five days, the animals are sacrificed and the Matrigel plugs, in which neovascularization can be visualized, are photographed. An experimental animal receiving Matrigel and an effective dose of test compound will exhibit

-87-

less vascularization than a control animal or an experimental animal receiving a less- or non-effective does of compound.

An *in vivo* system designed to test compounds for their ability to limit the spread of primary tumors is described by Crowley et al., *Proc. Natl. Acad. Sci.*, 90:5021-5025 (1993). Nude mice are injected with tumor cells (PC3) engineered to express CAT (chloramphenical acetyltransferase). Compounds to be tested for their ability to decrease tumor size and/or metastases are administered to the animals, and subsequent measurements of tumor size and/or metastatic growths are made. In addition, the level of CAT detected in various organs provides an indication of the ability of the test compound to inhibit metastasis; detection of less CAT in tissues of a treated animal versus a control animal indicates less CAT-expressing cells migrated to that tissue.

5

10

15

20

25

30

In vivo experimental modes designed to evaluate the inhibitory potential of a test serine protease inhibitors, using a tumor cell line F3II, the to be highly invasive, are described by Alonso et al., Breast Canc. Res. Treat., 40:209-223 (1996). This group describes in vivo studies for toxicity determination, tumor growth, invasiveness, spontaneous metastasis, experimental lung metastasis, and an angiogenesis assay.

The CAM model (chick embryo chorioallantoic membrane model), first described by L. Ossowski in 1998 (*J. Cell Biol.*, 107:2437-2445 (1988)), provides another method for evaluating the urokinase inhibitory activity of a test compound. In the CAM model, tumor cells invade through the chorioallantoic membrane containing CAM with tumor cells in the presence of several serine protease inhibitors results in less or no invasion of the tumor cells through the membrane. Thus, the CAM assay is performed with CAM and tumor cells in the presence and absence of various concentrations of test compound. The invasiveness of tumor cells is measured under such conditions to provide an indication of the compound's inhibitory activity. A compound having inhibitory activity correlates with less tumor invasion.

The CAM model is also used in a standard assay of angiogenesis (*i.e.*, effect on formation of new blood vessels (Brooks et al., *Methods in Molecular Biology*, 129:257-269 (1999)). According to this model, a filter disc containing

-88-

an angiogenesis inducer, such as basic fibroblast growth factor (bFDG) is placed onto the CAM. Diffusion of the cytokine into the CAM induces local angiogenesis, which may be measured in several ways such as by counting the number of blood vessel branch points within the CAM directly below the filter disc. The ability of identified compounds to inhibit cytokine-induced angiogenesis can be tested using this model. A test compound can either be added to the filter disc that contains the angiogenesis inducer, be placed directly on the membrane or be administered systemically. The extent of new blood vessel formation in the presence and/or absence of test compound can be compared using this model. The formation of fewer new blood vessels in the presence of a test compound would be indicative of anti-angiogenesis activity. Demonstration of anti-angiogenesis activity for inhibitors of an MTSP protein indicates a role in angiogenesis for that MTSP protein.

10

20

25

30

#### b. Known serine protease inhibitors

15 Compounds for screening can be serine protease inhibitors, which can be tested for their ability to inhibit the activity of an MTSP, particularly an MTSP3, MTSP4, or MTSP6.

Exemplary, but not limiting serine proteases, include the following known serine protease inhibitors are used in the screening assays: Serine Protease Inhibitor 3 (SPI-3) (Chen, M.C., et al., *Citokine*, 11(11):856-862 (1999)); Aprotinin (Iijima, R., et al., *J. Biochem. (Tokyo)*, 126(5):912-916 (1999)); Kazaltype serine protease inhibitor-like proteins (Niimi, T., et al., *Eur. J. Biochem.*, 266(1):282-292 (1999)); Kunitz-type serine protease inhibitor (Ravichandran, S., et al., *Acta Crystallogr. D. Biol. Crystallogr.*, 55(11):1814-1821 (1999)); Tissue factor pathway inhibitor-2/Matrix-associated serine rotease inhibitor (TFPI-2/MSPI), (Liu, Y., et al., *Arch. Biochem. Biophys.*, 370(1):112-8 (1999)); Bukunin, (Cui, C.Y., et al., *J. Invest. Dermatol.*, 113(2):182-8 (1999)); Nafmostat mesilate (Ryo, R., et al., *Vox Sang.*, 76(4):241-6 (1999)); TPCK (Huang, Y., et al., *Oncogene*, 18(23):3431-9 (1999)); A synthetic cotton-bound serine protease inhibitor (Edwards, J.V., et al., *Wound Repair Regen.*, 7(2):106-18 (1999)); FUT-175 (Sawada, M., et al., *Stroke*, 30(3):644-50 (1999)); Combination of serine protease inhibitor FUT-0175 and thromboxane synthetase

inhibitor OKY-046 (Kaminogo, M., et al., Neurol. Med. Chir. (Tokyo), 38(11):704-8; discussion 708-9 (1998)); The rat serine protease inhibitor 2.1 gene (LeCam, A., et al., Biochem. Biophys. Res. Commun., 253(2):311-4 (1998)); A new intracellular serine protease inhibitor expressed in the rat pituitary gland complexes with granzyme B (Hill, R.M., et al., FEBS Lett., 5 440(3):361-4 (1998)); 3,4-Dichloroisocoumarin (Hammed, A., et al., Proc. Soc. Exp. Biol. Med., 219(2):132-7 (1998)); LEXO32 (Bains, A.S., et al., Eur. J. Pharmacol., 356(1):67-72 (1998)); N-tosyl-L-phenylalanine chloromethyl ketone (Dryjanski, M., et al., Biochemistry, 37(40):14151-6 (1998)); Mouse gene for 10 the serine protease inhibitor neuroserpin (P112) (Berger, P., et al., Gene, 214(1-2):25-33 (1998)); Rat serine protease inhibitor 2.3 gene (Paul, C., et al., Eur. J. Biochem., 254(3):538-46 (1998)); Ecotin (Yang, S.Q., et al., J. Mol. Biol., 279(4):945-57 (1998)); A 14 kDa plant-related serine protease inhibitor (Roch, P., et al., Dev. Comp. Immunol., 22(1):1-12 (1998)); Matrix-associated serine 15 protease inhibitor TFPI-2/33 kDa MSPI (Rao, C.N., et al., Int. J. Cancer, 76(5):749-56 (1998)); ONO-3403 (Hiwasa, T., et al., Cancer Lett., 126(2):221-5 (1998)); Bdellastasin (Moser, M., et al., Eur. J. Biochem., 253(1):212-20 (1998)); Bikunin (Xu, Y., et al., J. Mol. Biol., 276(5):955-66 (1998)); Nafamostat mesilate (Mellgren, K., et al., Thromb. Haemost., 79(2):342-7 20 (1998)); The growth hormone dependent serine protease inhibitor, Spi 2.1 (Maake, C., et al., Endocrinology, 138(12):5630-6 (1997)); Growth factor activator inhibitor type 2, a Kunitz-type serine protease inhibitor (Kawaguchi, T., et al., J. Biol. Chem., 272(44):27558-64 (1997)); Heat-stable serine protease inhibitor protein from ovaries of the desert locust, Schistocerga gregaria 25 (Hamdaoui, A., et al., Biochem. Biophys. Res. Commun., 238(2):357-60 (1997)); Bikunin, (Delaria, K.A., et al., J. Biol. Chem., 272(18):12209-14 (1997)); Human placental bikunin (Marlor, C.W., et al., J. Biol. Chem., 272(10):12202-8 (1997)); Hepatocyte growth factor activator inhibitor, a novel Kunitz-type serine protease inhibitor (Shimomura, T., et al., J. Biol. Chem., 272(10):6370-6 (1997)); FUT-187, oral serine protease inhibitor, (Shiozaki, H., 30 et al., Gan To Kaguku Ryoho, 23(14): 1971-9 (1996)); Extracellular matrix-

associated serine protease inhibitors (Mr 33,000, 31,000, and 27,000 (Rao,

WO 01/57194

C.N., et al., Arch. Biochem. Biophys., 335(1):82-92 (1996)); An irreversible isocoumarin serine protease inhibitor (Palencia, D.D., et al., Biol. Reprod., 55(3):536-42 (1996)); 4-(2-aminoethyl)-benzenesulfonyl fluoride (AEBSF) (Nakabo, Y., et al., J. Leukoc. Biol., 60(3):328-36 (1996)); Neuroserpin (Osterwalder, T., et al., EMBO J., 15(12):2944-53 (1996)); Human serine 5 protease inhibitor alpha-1-antitrypsin (Forney, J.R., et al., J. Parasitol.. 82(3):496-502 (1996)); Rat serine protease inhibitor 2.3 (Simar-Blanchet, A.E., et al., Eur. J. Biochem., 236(2):638-48 (1996)); Gebaxate mesilate (parodi, F., et al., J. Cardiothorac. Vasc. Anesth., 10(2):235-7 (1996)); Recombinant serine protease inhibitor, CPTI II (Stankiewicz, M., et al., (Acta Biochim. Pol., 10 43(3):525-9 (1996)); A cysteine-rich serine protease inhibitor (Guamerin II) (Kim, D.R., et al., J. Enzym. Inhib., 10(2):81-91 (1996)); Diisopropylfluorophosphate (Lundqvist, H., et al., Inflamm. Res., 44(12):510-7 (1995)); Nexin 1 (Yu, D.W., et al., J. Cell Sci., 108(Pt 12):3867-74 (1995)); LEX032 (Scalia, R., et al., Shock, 4(4):251-6 (1995)); Protease nexin I (Houenou, L.J., et al., Proc. Natl. 15 Acad. Sci. U.S.A., 92(3):895-9 (1995)); Chymase-directed serine protease inhibitor (Woodard S.L., et al., J. Immunol., 153(11):5016-25 (1994)); N-alphatosyl-L-lysyl-chloromethyl ketone (TLCK) (Bourinbaiar, A.S., et al., Cell Immunol., 155(1):230-6 (1994)); Smpi56 (Ghendler, Y., et al., Exp. Parasitol., 78(2):121-31 (1994)); Schistosoma haematobium serine protease (Blanton, R.E., et al., 20 Mol. Biochem. Parasitol., 63(1):1-11 (1994)); Spi-1 (Warren, W.C., et al., Mol. Cell Endocrinol., 98(1):27-32 (1993)); TAME (Jessop, J.J., et al., Inflammation, 17(5):613-31 (1993)); Antithrombin III (Kalaria, R.N., et al., Am. J. Pathol., 143(3):886-93 (1993)); FOY-305 (Ohkoshi, M., et al., Anticancer Res., 25 13(4):963-6 (1993)); Camostat mesilate (Senda, S., et al., *Intern. Med.*, 32(4):350-4 (1993)); Pigment epithelium-derived factor (Steele, F.R., et al., Proc. Natl. Acad. Sci. U.S.A., 90(4):1526-30 (1993)); Antistasin (Holstein, T.W., et al., *FEBS Lett.*, 309(3):288-92 (1992)); The vaccinia virus K2L gene encodes a serine protease inhibitor (Zhou, J., et al., Virology, 189(2):678-86 30 (1992)); Bowman-Birk serine-protease inhibitor (Werner, M.H., et al., J. Mol. Biol., 225(3):873-89 (1992); FUT-175 (Yanamoto, H., et al., Neurosurgery,

30(3):358-63 (1992)); FUT-175; (Yanamoto, H., et al., Neurosurgery,

25

30

30(3):351-6, discussion 356-7 (1992)); PAI-I (Yreadwell, B.V., et al., J. Orthop. Res., 9(3):309-16 (1991)); 3,4-Dichloroisocoumarin (Rusbridge, N.M., et al., FEBS Lett., 268(1):133-6 (1990)); Alpha 1-antichymotrypsin (Lindmark, B.E., et al., Am. Rev. Respir. Des., 141(4 Pt 1):884-8 (1990)); P-toluenesulfonyl-Larginine methyl ester (TAME) (Scuderi, P., J. Immunol., 143(1):168-73 (1989)); 5 Aprotinin (Seto, S., et al., Adv. Exp. Med. Biol., 247B:49-54 (1989)); Alpha 1antichymotrypsin (Abraham, C.R., et al., Cell, 52(4):487-501 (1988)); Contrapsin (Modha, J., et al., Parasitology, 96 (Pt 1):99-109 (1988)); (FOY-305) (Yamauchi, Y., et al., Hiroshima J. Med. Sci., 36(1):81-7 No abstract available (1987)); Alpha 2-antiplasmin (Holmes, W.E., et al., J. Biol. Chem., 262(4):1659-10 64 (1987)); 3,4-dichloroisocoumarin (Harper, J.W., et al., Biochemistry, 24(8):1831-41 (1985)); Diisoprophylfluorophosphate (Tsutsui, K., et al., Biochem. Biophys. Res. Commun., 123(1):271-7 (1984)); Gabexate mesilate (Hesse, B., et al., Pharmacol. Res. Commun., 16(7):637-45 (1984)); Phenyl 15 methyl sulfonyl fluoride (Dufer, J., et al., Scand. J. Haematol., 32(1):25-32 (1984)); Aprotinin (Seto, S., et al., Hypertension, 5(6):893-9 (1983)); Protease inhibitor CI-2 (McPhalen, C.A., et al., J. Mol. Biol., 168(2):445-7 (1983)); Phenylmethylsulfonyl fluoride (Sekar V., et al., Biochem. Biophys. Res. Commun., 89(2):474-8 (1979)); PGE1 (Feinstein, M.D., et al., Prostaglandine, 20 14(6):1075-93 (1977)

## c. Combinatorial libraries and other libraries

The source of compounds for the screening assays, can be libraries, including, but are not limited to, combinatorial libraries. Methods for synthesizing combinatorial libraries and characteristics of such combinatorial libraries are known in the art (*See generally, Combinatorial Libraries: Synthesis, Screening and Application Potential* (Cortese Ed.) Walter de Gruyter, Inc., 1995; Tietze and Lieb, *Curr. Opin. Chem. Biol.*, 2(3):363-71 (1998); Lam, *Anticancer Drug Des.*, 12(3):145-67 (1997); Blaney and Martin, *Curr. Opin. Chem. Biol.*, 1(1):54-9 (1997); and Schultz and Schultz, *Biotechnol. Prog.*, 12(6):729-43 (1996)).

Methods and strategies for generating diverse libraries, primarily peptideand nucleotide-based oligomer libraries, have been developed using molecular

biology methods and/or simultaneous chemical synthesis methodologies (see, e.g., Dower et al., Annu. Rep. Med. Chem., 26:271-280 (1991); Fodor et al., Science, 251:767-773 (1991); Jung et al., Angew. Chem. Ind. Ed. Engl., 31:367-383 (1992); Zuckerman et al., Proc. Natl. Acad. Sci. USA, 89:4505-4509 (1992); Scott et al., Science, 249:386-390 (1990); Devlin et al., Science, 249:404-406 (1990); Cwirla et al., Proc. Natl. Acad. Sci. USA, 87:6378-6382 (1990); and Gallop et al., J. Medicinal Chemistry, 37:1233-1251 (1994)). The resulting combinatorial libraries potentially contain millions of compounds and that can be screened to identify compounds that exhibit a selected activity.

5

10

15

20

25

30

The libraries fall into roughly three categories: fusion-protein-displayed peptide libraries in which random peptides or proteins are presented on the surface of phage particles or proteins expressed from plasmids; support-bound synthetic chemical libraries in which individual compounds or mixtures of compounds are presented on insoluble matrices, such as resin beads (see, e.g., Lam et al., Nature, 354:82-84 (1991)) and cotton supports (see, e.g., Eichler et al., Biochemistry 32:11035-11041 (1993)); and methods in which the compounds are used in solution (see, e.g., Houghten et al., Nature, 354:84-86 (1991); Houghten et al., BioTechniques, 313:412-421 (1992); and Scott et al., Curr. Opin. Biotechnol., 5:40-48 (1994)). There are numerous examples of synthetic peptide and oligonucleotide combinatorial libraries and there are many methods for producing libraries that contain non-peptidic small organic molecules. Such libraries can be based on basis set of monomers that are combined to form mixtures of diverse organic molecules or that can be combined to form a library based upon a selected pharmacophore monomer.

Either a random or a deterministic combinatorial library can be screened by the presently disclosed and/or claimed screening methods. In either of these two libraries, each unit of the library is isolated and/or immobilized on a solid support. In the deterministic library, one knows a priori a particular unit's location on each solid support. In a random library, the location of a particular unit is not known a priori although each site still contains a single unique unit. Many methods for preparing libraries are known to those of skill in this art (see, e.g., Geysen et al., Proc. Natl. Acad. Sci. USA, 81:3998-4002 (1984),

Houghten et al., Proc. Natl. Acad. Sci. USA, 81:5131-5135 (1985)). Combinatorial library generated by the any techniques known to those of skill in the art are contemplated (see, e.g., Table 1 of Schultz and Schultz, Biotechnol. Prog., 12(6):729-43 (1996)) for screening; Bartel et al., Science, 261:1411-1418 (1993); Baumbach et al. BioPharm, (May):24-35 (1992); Bock et al. 5 Nature, 355:564-566 (1992); Borman, S., Combinatorial chemists focus on small molecules molecular recognition, and automation, Chem. Eng. News, 2(12):29 (1996); Boublik, et al., Eukaryotic Virus Display: Engineering the Major Surface Glycoproteins of the Autographa California Nuclear Polyhedrosis Virus 10 (ACNPV) for the Presentation of Foreign Proteins on the Virus Surface, Bio/Technology, 13:1079-1084 (1995); Brenner, et al., Encoded Combinatorial Chemistry, Proc. Natl. Acad Sci. U.S.A., 89:5381-5383 (1992); Caflisch, et al., Computational Combinatorial Chemistry for De Novo Ligand Design: Review and Assessment, Perspect. Drug Discovery Des., 3:51-84 (1995); Cheng, et al., Sequence-Selective Peptide Binding with a Peptido-A,B-trans-steroidal Receptor 15 Selected from an Encoded Combinatorial Library, J. Am. Chem. Soc., 118:1813-1814 (1996); Chu, et al., Affinity Capillary Electrophoresis to Identify the Peptide in A Peptide Library that Binds Most Tightly to Vancomycin, J. Org. Chem., 58:648-652 (1993); Clackson, et al., Making Antibody Fragments Using Phage Display Libraries, Nature, 352:624-628 (1991); Combs, et al., Protein 20 Structure-Based Combinatorial Chemistry: Discovery of Non-Peptide Binding Elements to Src SH3 Domain, J. Am. Chem. Soc., 118:287-288 (1996); Cwirla, et al., Peptides On Phage: A Vast Library of Peptides for Identifying Ligands, Proc. Natl. Acad. Sci. U.S.A., 87:6378-6382 (1990); Ecker, et al., Combinatorial 25 Drug Discovery: Which Method will Produce the Greatest Value, Bio/Technology, 13:351-360 (1995); Ellington, et al., In Vitro Selection of RNA Molecules That Bind Specific Ligands, Nature, 346:818-822 (1990); Ellman, J.A., Variants of Benzodiazephines, J. Am. Chem. Soc., <u>114</u>:10997 (1992); Erickson, et al., The Proteins; Neurath, H., Hill, R.L., Eds.: Academic: New York, 30 1976; pp. 255-257; Felici, et al., J. Mol. Biol., 222:301-310 (1991); Fodor, et al., Light-Directed, Spatially Addressable Parallel Chemical Synthesis, Science,

251:767-773 (1991); Francisco, et al., Transport and Anchoring of Beta-

Lactamase to the External Surface of E. Coli., Proc. Natl. Acad. Sci. U.S.A., 89:2713-2717 (1992); Georgiou, et al., Practical Applications of Engineering Gram-Negative Bacterial Cell Surfaces, TIBTECH, 11:6-10 (1993); Geysen, et al., Use of peptide synthesis to probe viral antigens for epitopes to a resolution of a single amino acid, Proc. Natl. Acad. Sci. U.S.A., 81:3998-4002 (1984); Glaser, 5 et al., Antibody Engineering by Condon-Based Mutagenesis in a Filamentous Phage Vector System, J. Immunol., 149:3903-3913 (1992); Gram, et al., In vitro selection and affinity maturation of antibodies from a naive combinatorial immunoglobulin library, Proc. Natl. Acad. Sci., 89:3576-3580 (1992); Han, et al., Liquid-Phase Combinatorial Synthesis, Proc. Natl. Acad. Sci. U.S.A., 10 92:6419-6423 (1995); Hoogenboom, et al., Multi-Subunit Proteins on the Surface of Filamentous Phage: Methodologies for Displaying Antibody (Fab) Heavy and Light Chains, Nucleic Acids Res., 19:4133-4137 (1991); Houghten, et al., General Method for the Rapid Solid-Phase Synthesis of Large Numbers of Peptides: Specificity of Antigen-Antibody Interaction at the Level of Individual 15 Amino Acids, Proc. Natl. Acad. Sci. U.S.A., 82:5131-5135 (1985); Houghten, et al., The Use of Synthetic Peptide Combinatorial Libraries for the Determination of Peptide Ligands in Radio-Receptor Assays-Opiod-Peptides, Bioorg. Med. . Chem. Lett., 3:405-412 (1993); Houghten, et al., Generation and Use of 20 Synthetic Peptide Combinatorial Libraries for Basic Research and Drug Discovery, Nature, 354:84-86 (1991); Huang, et al., Discovery of New Ligand Binding Pathways in Myoglobin by Random Mutagenesis, Nature Struct. Biol., 1:226-229 (1994); Huse, et al., Generation of a Large Combinatorial Library of the Immunoglobulin Repertoire In Phage Lambda, Science, 246:1275-1281 (1989); Janda, K.D., New Strategies for the Design of Catalytic Antibodies, Biotechnol. 25 Prog., 6:178-181 (1990); Jung, et al., Multiple Peptide Synthesis Methods and Their Applications, Angew. Chem. Int. Ed. Engl., 31:367-486 (1992); Kang, et al., Linkage of Recognition and Replication Functions By Assembling Combinatorial Antibody Fab Libraries Along Phage Surfaces, Proc. Natl. Acad. Sci. U.S.A., 88:4363-4366 (1991a); Kang, et al., Antibody Redesign by Chain 30 Shuffling from Random Combinatorial Immunoglobulin Libraries, Proc. Natl. Acad. Sci. U.S.A., 88:11120-11123 (1991b); Kay, et al., An M13 Phage Library

WO 01/57194

Displaying Random 38-Amino-Acid-Peptides as a Source of Novel Sequences with Affinity to Selected Targets Genes, Gene, 128:59-65 (1993); Lam, et al., A new type of synthetic peptide library for identifying ligand-binding activity, Nature, 354:82-84 (1991) (published errata apear in Nature, 358:434 (1992) and Nature, 360:768 (1992); Lebl, et al., One Bead One Structure Combinatorial 5 Libraries, Biopolymers (Pept. Sci.), 37:177-198 (1995); Lerner, et al., Antibodies without Immunization, Science, 258:1313-1314 (1992); Li, et al., Minimization of a Polypeptide Hormone, Science, 270:1657-1660 (1995); Light, et al., Display of Dimeric Bacterial Alkaline Phosphatase on the Major Coat Protein of Filamentous Bacteriophage, Bioorg. Med. Chem. Lett., 3:1073-1079 (1992); 10 Little, et al., Bacterial Surface Presentation of Proteins and Peptides: An Alternative to Phage Technology, Trends Biotechnol., 11:3-5 (1993); Marks, et al., By-Passing Immunization. Human Antibodies from V-Gene Libraries Displayed on Phage, J. Mol. Biol., 222:581-597 (1991); Matthews, et al., Substrate Phage: Selection of Protease Substrates by Monovalent Phage Display, 15 Science, 260:1113-1117 (1993); McCafferty, et al., Phage Enzymes: Expression and Affinity Chromatography of Functional Alkaline Phosphatase on the Surface of Bacteriophage, Protein Eng., 4:955-961 (1991); Menger, et al., Phosphatase Catalysis Developed Via Combinatorial Organic Chemistry, J. Org. Chem., 20 60:6666-6667 (1995); Nicolaou, et al., Angew. Chem. Int. Ed. Engl., 34:2289-2291 (1995); Oldenburg, et al., Peptide Ligands for A Sugar-Binding Protein Isolated from a Random Peptide Library, Proc. Natl. Acad. Sci. U.S.A., 89:5393-5397 (1992); Parmley, et al., Antibody-Selectable Filamentous fd Phage Vectors: Affinity Purification of Target Genes, Genes, 73:305-318 (1988); Pinilla, et al., Synthetic Peptide Combinatorial Libraries (SPCLS)--Identification of the Antigenic 25 Determinant of Beta-Endorphin Recognized by Monoclonal Antibody-3E7, Gene, 128:71-76 (1993); Pinilla, et al., Review of the Utility of Soluble Combinatorial Libraries, Biopolymers, 37:221-240 (1995); Pistor, et al., Expression of Viral Hemegglutinan On the Surface of E. Coli., Klin. Wochenschr., 66:110-116 (1989); Pollack, et al., Selective Chemical Catalysis by an Antibody, Science, 30 234:1570-1572 (1986); Rigler, et al., Fluorescence Correlations, Single Molecule Detection and Large Number Screening: Applications in Biotechnology, J.

-96-

Biotechnol., 41:177-186 (1995); Sarvetnick, et al., Increasing the Chemical Potential of the Germ-Line Antibody Repertoire, Proc. Natl. Acad. Sci. U.S.A., 90:4008-4011 (1993); Sastry, et al., Cloning of the Immunological Repertiore in Escherichia Coli for Generation of Monoclonal Catalytic Antibodies: Construction of a Heavy Chain Variable Region-Specific cDNA Library, Proc. Natl. Acad. Sci. 5 U.S.A., 86:5728-5732 (1989); Scott, et al., Searching for Peptide Ligands with an Epitope Library, Science, 249:386-390 (1990); Sears, et al., Engineering Enzymes for Bioorganic Synthesis: Peptide Bond Formation, Biotechnol. Prog., 12:423-433 (1996); Simon, et. al., Peptides: A Modular Approach to Drug Discovery, Proc. Natl. Acad. Sci. U.S.A., 89:9367-9371 (1992); Still, et al., 10 Discovery of Sequence-Selective Peptide Binding by Synthetic Receptors Using Encoded Combinatorial Libraries, Acc. Chem. Res., 29:155-163 (1996); Thompson, et al., Synthesis and Applications of Small Molecule Libraries, Chem. Rev., 96:555-600 (1996); Tramontano, et al., Catalytic Antibodies, Science, 234:1566-1570 (1986); Wrighton, et al., Small Peptides as Potent Mimetics of 15 the Protein Hormone Erythropoietin, Science, 273:458-464 (1996); York, et al., Combinatorial mutagenesis of the reactive site region in plasminogen activator inhibitor I. J. Biol. Chem., 266:8595-8600 (1991); Zebedee, et al., Human Combinatorial Antibody Libraries to Hepatitis B Surface Antigen, Proc. Natl. Acad. Sci. U.S.A., 89:3175-3179 (1992); Zuckerman, et al., Identification of 20 Highest-Affinity Ligands by Affinity Selection from Equimolar Peptide Mixtures Generated by Robotic Synthesis, Proc. Natl. Acad. Sci. U.S.A., 89:4505-4509 (1992).

For example, peptides that bind to an MTSP protein or a protease domain of an MTSP protein can be identified using phage display libraries. In an exemplary embodiment, this method can include a) contacting phage from a phage library with the MTSP protein or a protease domain thereof; (b) isolating phage that bind to the protein; and (c) determining the identity of at least one peptide coded by the isolated phage to identify a peptide that binds to an MTSP protein.

25

30

#### H. Modulators of the activity of MTSP proteins

Provided herein are compounds, identified by screening or produced using the MTSP proteins or protease domain in other screening methods, that modulate the activity of an MTSP. These compounds act by directly interacting with the MTSP protein or by altering transcription or translation thereof. Such molecules include, but are not limited to, antibodies that specifically react with an MTSP protein, particularly with the protease domain thereof, antisense nucleic acids that alter expression of the MTSP protein, antibodies, peptide mimetics and other such compounds.

#### 1. Antibodies

5

10

15

20

25

30

Antibodies, including polyclonal and monoclonal antibodies, that specifically bind to the MTSP proteins provided herein, particularly to the single chain protease domains thereof are provided. Preferably, the antibody is a monoclonal antibody, and preferably, the antibody specifically binds to the protease domain of the MTSP protein. In particular embodiments, antibodies to each of the single chain of protease domain of MTSP1, MTSP3, MTSP4 and MTSP6. Also provided are antibodies that specifically bind to any domain of MTSP3 or MTSP4, and to double chain forms thereof.

The MTSP protein and domains, fragments, homologs and derivatives thereof may be used as immunogens to generate antibodies that specifically bind such immunogens. Such antibodies include but are not limited to polyclonal, monoclonal, chimeric, single chain, Fab fragments, and an Fab expression library. In a specific embodiment, antibodies to human MTSP protein are produced. In another embodiment, complexes formed from fragments of MTSP protein, which fragments contain the serine protease domain, are used as immunogens for antibody production.

Various procedures known in the art may be used for the production of polyclonal antibodies to MTSP protein, its domains, derivatives, fragments or analogs. For production of the antibody, various host animals can be immunized by injection with the native MTSP protein or a synthetic version, or a derivative of the foregoing, such as a cross-linked MTSP protein. Such host animals include but are not limited to rabbits, mice, rats, etc. Various adjuvants can be

used to increase the immunological response, depending on the host species, and include but are not limited to Freund's (complete and incomplete), mineral gels such as aluminum hydroxide, surface active substances such as lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, dinitrophenol, and potentially useful human adjuvants such as bacille Calmette-Guerin (BCG) and corynebacterium parvum.

5

10

15

20

25

30

For preparation of monoclonal antibodies directed towards an MTSP protein or domains, derivatives, fragments or analogs thereof, any technique that provides for the production of antibody molecules by continuous cell lines in culture may be used. Such techniques include but are not restricted to the hybridoma technique originally developed by Kohler and Milstein (Nature 256:495-497 (1975)), the trioma technique, the human B-cell hybridoma technique (Kozbor et al., Immunology Today 4:72 (1983)), and the EBV hybridoma technique to produce human monoclonal antibodies (Cole et al., in Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, Inc., pp. 77-96 (1985)). In an additional embodiment, monoclonal antibodies can be produced in germ-free animals utilizing recent technology (PCT/US90/02545). Human antibodies may be used and can be obtained by using human hybridomas (Cote et al., Proc. Natl. Acad. Sci. USA 80:2026-2030 (1983)). Or by transforming human B cells with EBV virus in vitro (Cole et al., in Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, Inc., pp. 77-96 (1985)). Techniques developed for the production of "chimeric antibodies" (Morrison et al., Proc. Natl. Acad. Sci. USA 81:6851-6855 (1984); Neuberger et al., Nature 312:604-608 (1984); Takeda et al., Nature 314:452-454 (1985)) by splicing the genes from a mouse antibody molecule specific for the MTSP protein together with genes from a human antibody molecule of appropriate biological activity can be used.

Techniques described for the production of single chain antibodies (U.S. patent 4,946,778) can be adapted to produce MTSP protein-specific single chain antibodies. An additional embodiment uses the techniques described for the construction of Fab expression libraries (Huse et al., *Science* 246:1275-1281 (1989)) to allow rapid and easy identification of monoclonal Fab fragments with the desired specificity for MTSP protein or MTSP protein, or domains,

-99-

derivatives, or analogs thereof. Non-human antibodies can be "humanized" by known methods (see, e.g., U.S. Patent No. 5,225,539).

Antibody fragments that contain the idiotypes of MTSP protein can be generated by techniques known in the art. For example, such fragments include but are not limited to: the F(ab')2 fragment which can be produced by pepsin digestion of the antibody molecule; the Fab' fragments that can be generated by reducing the disulfide bridges of the F(ab')2 fragment, the Fab fragments that can be generated by treating the antibody molecular with papain and a reducing agent, and Fv fragments.

In the production of antibodies, screening for the desired antibody can be accomplished by techniques known in the art, e.g., ELISA (enzyme-linked immunosorbent assay). To select antibodies specific to a particular domain of the MTSP protein one may assay generated hybridomas for a product that binds to the fragment of the MTSP protein that contains such a domain

The foregoing antibodies can be used in methods known in the art relating to the localization and/or quantitation of MTSP proteins, *e.g.*, for imaging these proteins, measuring levels thereof in appropriate physiological samples, in diagnostic methods, etc.

In another embodiment, (see *infra*), anti-MTSP protein antibodies, or fragments thereof, containing the binding domain are used as therapeutic agents.

### 2. Peptides and Peptide Mimetics

5

10

15

20

25

30

Provided herein are methods for identifying molecules that bind to and modulate the activity of MTSP proteins. Included among molecules that bind to MTSPs, particularly the single chain protease domain or catalytically active fragments thereof, are peptides and peptide mimetics. Peptide mimetics are molecules or compounds that mimic the necessary molecular conformation of a ligand or polypeptide for specific binding to a target molecule such as, *e.g.*, an MTSP protein. In an exemplary embodiment, the peptides or peptide mimetics bind to the protease domain of the MTSP protein. Such peptides and peptide mimetics include those of antibodies that specifically bind an MTSP protein and, preferably, bind to the protease domain of an MTSP protein. The peptides and

-100-

peptide mimetics identified by methods provided herein can be agonists or antagonists of MTSP proteins.

Such peptides and peptide mimetics are useful for diagnosing, treating, preventing, and screening for a disease or disorder associated with MTSP protein activity in a mammal. In addition, the peptides and peptide mimetics are useful for identifying, isolating, and purifying molecules or compounds that modulate the activity of an MTSP protein, or specifically bind to an MTSP protein, preferably, the protease domain of an MTSP protein. Low molecular weight peptides and peptide mimetics can have strong binding properties to a target molecule, *e.g.*, an MTSP protein or, preferably, to the protease domain of an MTSP protein.

10

15

20

25

30

Peptides and peptide mimetics that bind to MTSP proteins as described herein can be administered to mammals, including humans, to modulate MTSP protein activity. Thus, methods for therapeutic treatment and prevention of neoplastic diseases comprise administering a peptide or peptide mimetic compound in an amount sufficient to modulate such activity are provided. Thus, also provided herein are methods for treating a subject having such a disease or disorder in which a peptide or peptide mimetic compound is administered to the subject in a therapeutically effective dose or amount.

Compositions containing the peptides or peptide mimetics can be administered for prophylactic and/or therapeutic treatments. In therapeutic applications, compositions can be administered to a patient already suffering from a disease, as described above, in an amount sufficient to cure or at least partially arrest the symptoms of the disease and its complications. Amounts effective for this use will depend on the severity of the disease and the weight and general state of the patient.

In prophylactic applications, compositions containing the peptides and peptide mimetics are administered to a patient susceptible to or otherwise at risk of a particular disease. Such an amount is defined to be a "prophylactically effective dose". In this use, the precise amounts again depend on the patient's state of health and weight.

-101-

Accordingly, the peptides and peptide mimetics that bind to an MTSP protein can be used generating pharmaceutical compositions containing, as an active ingredient, at least one of the peptides or peptide mimetics in association with a pharmaceutical carrier or diluent. The compounds can be administered, for example, by oral, pulmonary, parental (intramuscular, intraperitoneal, intravenous (IV) or subcutaneous injection), inhalation (via a fine powder formulation), transdermal, nasal, vaginal, rectal, or sublingual routes of administration and can be formulated in dosage forms appropriate for each route of administration (see, e.g., International PCT application Nos. WO 93/25221 and WO 94/17784; and European Patent Application 613,683).

10

15

20

25

30

Peptides and peptide mimetics that bind to MTSP proteins are useful *in vitro* as unique tools for understanding the biological role of MTSP proteins, including the evaluation of the many factors thought to influence, and be influenced by, the production of MTSP protein. Such peptides and peptide mimetics are also useful in the development of other compounds that bind to and modulate the activity of an MTSP protein, because such compounds provide important information on the relationship between structure and activity that should facilitate such development.

The peptides and peptide mimetics are also useful as competitive binders in assays to screen for new MTSP proteins or MTSP protein agonists. In such assay embodiments, the compounds can be used without modification or can be modified in a variety of ways; for example, by labeling, such as covalently or non-covalently joining a moiety which directly or indirectly provides a detectable signal. In any of these assays, the materials thereto can be labeled either directly or indirectly. Possibilities for direct labeling include label groups such as: radiolabels such as <sup>125</sup>I enzymes (U.S. Pat. No. 3,645,090) such as peroxidase and alkaline phosphatase, and fluorescent labels (U.S. Pat. No. 3,940,475) capable of monitoring the change in fluorescence intensity, wavelength shift, or fluorescence polarization. Possibilities for indirect labeling include biotinylation of one constituent followed by binding to avidin coupled to one of the above label groups. The compounds may also include spacers or linkers in cases where the compounds are to be attached to a solid support.

-102-

Moreover, based on their ability to bind to an MTSP protein, the peptides and peptide mimetics can be used as reagents for detecting MTSP proteins in living cells, fixed cells, in biological fluids, in tissue homogenates, in purified, natural biological materials, etc. For example, by labelling such peptides and peptide mimetics, one can identify cells having MTSP proteins. In addition, based on their ability to bind an MTSP protein, the peptides and peptide mimetics can be used in situ staining, FACS (fluorescence-activated cell sorting), Western blotting, ELISA, etc. In addition, based on their ability to bind to an MTSP protein, the peptides and peptide mimetics can be used in purification of MTSP protein polypeptides or in purifying cells expressing the MTSP protein polypeptides, e.g., a polypeptide encoding the protease domain of an MTSP protein.

The peptides and peptide mimetics can also be used as commercial reagents for various medical research and diagnostic uses.

The activity of the peptides and peptide mimetics can be evaluated either in vitro or in vivo in one of the numerous models described in McDonald (1992) Am. J. of Pediatric Hematology/Oncology, 14:8-21, which is incorporated herein by reference.

#### 3. Peptide and peptide mimetic therapy

5

10

15

20

Peptides and peptide mimetics that can bind to MTSP proteins or the protease domain of MTSP proteins and modulate the activity thereof, or have MTSP protein activity, can be used for treatment of neoplastic diseases. The peptides and peptide mimetics may be delivered, in vivo or ex vivo, to the cells of a subject in need of treatment. Further, peptides which have MTSP protein 25 activity can be delivered, in vivo or ex vivo, to cells which carry mutant or missing alleles encoding the MTSP protein gene. Any of the techniques described herein or known to the skilled artisan can be used for preparation and in vivo or ex vivo delivery of such peptides and peptide mimetics that are substantially free of other human proteins. For example, the peptides can be 30 readily prepared by expression in a microorganism or synthesis in vitro.

The peptides or peptide mimetics can be introduced into cells, in vivo or ex vivo, by microinjection or by use of liposomes, for example. Alternatively, the

-103-

peptides or peptide mimetics may be taken up by cells, in vivo or ex vivo, actively or by diffusion. In addition, extracellular application of the peptide or peptide mimetic may be sufficient to effect treatment of a neoplastic disease. Other molecules, such as drugs or organic compounds, that: 1) bind to an MTSP protein or protease domain thereof; or 2) have a similar function or activity to an MTSP protein or protease domain thereof, may be used in methods for treatment.

#### 4. Rational drug design

10

15

25

30

The goal of rational drug design is to produce structural analogs of biologically active polypeptides or peptides of interest or of small molecules or peptide mimetics with which they interact (e.g., agonists, antagonists, inhibitors) in order to fashion drugs which are, e.g., more active or stable forms thereof; or which, e.g., enhance or interfere with the function of a polypeptide in vivo (e.g., an MTSP protein). In one approach, one first determines the three-dimensional structure of a protein of interest (e.g., an MTSP protein or polypeptide having a protease domain) or, for example, of a MTSP protein-ligand complex, by X-ray crystallography, by computer modeling or most typically, by a combination of approaches (see, e.g., Erickson et al. 1990). Also, useful information regarding the structure of a polypeptide may be gained by modeling based on the structure 20 of homologous proteins. In addition, peptides can be analyzed by an alanine scan. In this technique, an amino acid residue is replaced by Ala, and its effect on the peptide's activity is determined. Each of the amino acid residues of the peptide is analyzed in this manner to determine the important regions of the peptide.

Also, a polypeptide or peptide that binds to an MTSP protein or, preferably, the protease domain of an MTSP protein, can be selected by a functional assay, and then the crystal structure of this polypeptide or peptide can be determined. The polypeptide can be, for example, an antibody specific for an MTSP protein or the protein domain of an MTSP protein. This approach can yield a pharmacore upon which subsequent drug design can be based. Further, it is possible to bypass the crystallography altogether by generating anti-idiotypic polypeptides or peptides, (anti-ids) to a functional,

-104-

pharmacologically active polypeptide or peptide that binds to an MTSP protein or protease domain of an MTSP protein. As a mirror image of a mirror image, the binding site of the anti-ids is expected to be an analog of the original target molecule, e.g., an MTSP protein or polypeptide having an MTSP protein. The anti-id could then be used to identify and isolate peptides from banks of chemically or biologically produced banks of peptides. Selected peptides would then act as the pharmacore.

5

10

15

20

25

30

Thus, one may design drugs which have, e.g., improved activity or stability or which act as modulators (e.g., inhibitors, agonists, antagonists, etc.) of MTSP protein activity, and are useful in the methods, particularly the methods for diagnosis, treatment, prevention, and screening of a neoplastic disease. By virtue of the availability of cloned MTSP protein sequences, sufficient amounts of the MTSP protein polypeptide may be made available to perform such analytical studies as X-ray crystallography. In addition, the knowledge of the amino acid sequence of an MTSP protein or the protease domain thereof, e.g., the protease domain encoded by the amino acid sequence of SEQ ID NO: 2, can provide guidance on computer modeling techniques in place of, or in addition to, X-ray crystallography.

# Methods of identifying peptides and peptide mimetics that bind to MTSP proteins

Peptides having a binding affinity to the MTSP protein polypeptides provided herein (e.g., an MTSP protein or a polypeptide having a protease domain of an MTSP protein) can be readily identified, for example, by random peptide diversity generating systems coupled with an affinity enrichment process. Specifically, random peptide diversity generating systems include the "peptides on plasmids" system (see, e.g., U.S. Patent Nos. 5,270,170 and 5,338,665); the "peptides on phage" system (see, e.g., U.S. Patent No. 6,121,238 and Cwirla,et al. (1990) Proc. Natl. Acad. Sci. U.S.A. 87:6378-6382); the "polysome system;" the "encoded synthetic library (ESL)" system; and the "very large scale immobilized polymer synthesis" system (see, e.g., U.S. Patent No. 6,121,238; and Dower et al. (1991) An. Rep. Med. Chem. 26:271-280

-105-

For example, using the procedures described above, random peptides can generally be designed to have a defined number of amino acid residues in length (e.g., 12). To generate the collection of oligonucleotides encoding the random peptides, the codon motif (NNK)x, where N is nucleotide A, C, G, or T (equimolar; depending on the methodology employed, other nucleotides can be employed), K is G or T (equimolar), and x is an integer corresponding to the number of amino acids in the peptide (e.g., 12) can be used to specify any one of the 32 possible codons resulting from the NNK motif: 1 for each of 12 amino acids, 2 for each of 5 amino acids, 3 for each of 3 amino acids, and only one of the three stop codons. Thus, the NNK motif encodes all of the amino acids, encodes only one stop codon, and reduces codon bias.

5

10

15

20

25

30

The random peptides can be presented, for example, either on the surface of a phage particle, as part of a fusion protein containing either the pIII or the pVIII coat protein of a phage fd derivative (peptides on phage) or as a fusion protein with the LacI peptide fusion protein bound to a plasmid (peptides on plasmids). The phage or plasmids, including the DNA encoding the peptides, can be identified and isolated by an affinity enrichment process using immobilized MTSP protein polypeptide having a protease domain. The affinity enrichment process, sometimes called "panning," typically involves multiple rounds of incubating the phage, plasmids, or polysomes with the immobilized MTSP protein polypeptide, collecting the phage, plasmids, or polysomes that bind to the MTSP protein polypeptide (along with the accompanying DNA or mRNA), and producing more of the phage or plasmids (along with the accompanying LacI-peptide fusion protein) collected.

#### Characteristics of peptides and peptide mimetics

Typically, the molecular weight of preferred peptides or peptide mimetics is from about 250 to about 8,000 daltons. If the peptides are oligomerized, dimerized and/or derivatized with a hydrophilic polymer (e.g., to increase the affinity and/or activity of the compounds), the molecular weights of such peptides can be substantially greater and can range anywhere from about 500 to about 120,000 daltons, more preferably from about 8,000 to about 80,000 daltons. Such peptides can comprise 9 or more amino acids wherein the amino

-106-

acids are naturally occurring or synthetic (non-naturally occurring) amino acids. One skilled in the art would know how to determine the affinity and molecular weight of the peptides and peptide mimetics suitable for therapeutic and/or diagnostic purposes (e.g., see Dower et al., U.S. Patent No. 6,121,238).

5

10

15

20

25

30

The peptides may be covalently attached to one or more of a variety of hydrophilic polymers. Suitable hydrophilic polymers include, but are not limited to, polyalkylethers as exemplified by polyethylene glycol and polypropylene glycol, polylactic acid, polyglycolic acid, polyoxyalkenes, polyvinylalcohol, polyvinylpyrrolidone, cellulose and cellulose derivatives, dextran and dextran derivatives, etc. When the peptide compounds are derivatized with such polymers, their solubility and circulation half-lives can be increased with little, if any, diminishment in their binding activity. The peptide compounds may be dimerized and each of the dimeric subunits can be covalently attached to a hydrophilic polymer. The peptide compounds can be PEGylated, i.e., covalently attached to polyethylene glycol (PEG).

Peptide analogs are commonly used in the pharmaceutical industry as non-peptide drugs with properties analogous to those of the template peptide. These types of non-peptide compounds are termed "peptide mimetics" or "peptidomimetics" (Luthman et al., A Textbook of Drug Design and Development, 14:386-406, 2nd Ed., Harwood Academic Publishers (1996); Joachim Grante (1994) Angew. Chem. Int. Ed. Engl., 33:1699-1720; Fauchere (1986) J. Adv. Drug Res., 15:29; Veber and Freidinger (1985) TINS, p. 392; and Evans et al. (1987) J. Med. Chem. 30:1229). Peptide mimetics that are structurally similar to therapeutically useful peptides may be used to produce an equivalent or enhanced therapeutic or prophylactic effect. Preparation of peptidomimetics and structures thereof are known to those of skill in this art.

Systematic substitution of one or more amino acids of a consensus sequence with a D-amino acid of the same type (e.g., D-lysine in place of L-lysine) may be used to generate more stable peptides. In addition, constrained peptides containing a consensus sequence or a substantially identical consensus sequence variation may be generated by methods known in the art (Rizo et al. (1992) An. Rev. Biochem., 61:387, incorporated herein by reference); for

-107-

example, by adding internal cysteine residues capable of forming intramolecular disulfide bridges which cyclize the peptide.

Those skilled in the art would appreciate that modifications may be made to the peptides and mimetics without deleteriously effecting the biological or functional activity of the peptide. Further, the skilled artisan would know how to design non-peptide structures in three dimensional terms, that mimic the peptides that bind to a target molecule, *e.g.*, an MTSP protein or, preferably, the protease domain of MTSP proteins (see, *e.g.*, Eck and Sprang (1989) *J. Biol. Chem.*, 26: 17605-18795).

10

15

20

25

30

When used for diagnostic purposes, the peptides and peptide mimetics may be labeled with a detectable label and, accordingly, the peptides and peptide mimetics without such a label can serve as intermediates in the preparation of labeled peptides and peptide mimetics. Detectable labels can be molecules or compounds, which when covalently attached to the peptides and peptide mimetics, permit detection of the peptide and peptide mimetics in vivo, for example, in a patient to whom the peptide or peptide mimetic has been administered, or in vitro, e.g., in a sample or cells. Suitable detectable labels are well known in the art and include, by way of example, radioisotopes, fluorescent labels (e.g., fluorescein), and the like. The particular detectable label employed is not critical and is selected relative to the amount of label to be employed as well as the toxicity of the label at the amount of label employed. Selection of the label relative to such factors is well within the skill of the art.

Covalent attachment of a detectable label to the peptide or peptide mimetic is accomplished by conventional methods well known in the art. For example, when the <sup>125</sup>I radioisotope is employed as the detectable label, covalent attachment of <sup>125</sup>I to the peptide or the peptide mimetic can be achieved by incorporating the amino acid tyrosine into the peptide or peptide mimetic and then iodinating the peptide (see, *e.g.*, Weaner *et al.* (1994) *Synthesis and Applications of Isotopically Labelled Compounds*, pp. 137-140). If tyrosine is not present in the peptide or peptide mimetic, incorporation of tyrosine to the N or C terminus of the peptide or peptide mimetic can be achieved by well known chemistry. Likewise, <sup>32</sup>P can be incorporated onto the peptide or peptide

-108-

mimetic as a phosphate moiety through, for example, a hydroxyl group on the peptide or peptide mimetic using conventional chemistry.

Labeling of peptidomimetics usually involves covalent attachment of one or more labels, directly or through a spacer (e.g., an amide group), to non-interfering position(s) on the peptidomimetic that are predicted by quantitative structure-activity data and/or molecular modeling. Such non-interfering positions generally are positions that do not form direct contacts with the macromolecules(s) to which the peptidomimetic binds to produce the therapeutic effect. Derivatization (e.g., labeling) of peptidomimetics should not substantially interfere with the desired biological or pharmacological activity of the peptidomimetic.

5

10

15

20

25

30

# 6. Methods of preparing peptides and peptide mimetics

Peptides that bind to MTSP proteins can be prepared by classical methods known in the art, for example, by using standard solid phase techniques. The standard methods include exclusive solid phase synthesis, partial solid phase synthesis methods, fragment condensation, classical solution synthesis, and even by recombinant DNA technology (see, *e.g.*, Merrifield (1963) *J. Am. Chem. Soc.*, 85:2149, incorporated herein by reference.)

Using the "encoded synthetic library" or "very large scale immobilized polymer synthesis" systems (see, e.g., U.S. Patent No. 5,925,525, and 5,902,723); one can not only determine the minimum size of a peptide with the activity of interest, one can also make all of the peptides that form the group of peptides that differ from the preferred motif (or the minimum size of that motif) in one, two, or more residues. This collection of peptides can then be screened for ability to bind to the target molecule, e.g., and MTSP protein or, preferably, the protease domain of an MTSP protein. This immobilized polymer synthesis system or other peptide synthesis methods can also be used to synthesize truncation analogs and deletion analogs and combinations of truncation and deletion analogs of the peptide compounds.

These procedures can also be used to synthesize peptides in which amino acids other than the 20 naturally occurring, genetically encoded amino acids are substituted at one, two, or more positions of the peptide. For instance,

naphthylalanine can be substituted for tryptophan, facilitating synthesis. Other synthetic amino acids that can be substituted into the peptides include L-hydroxypropyl, L-3, 4-dihydroxy-phenylalanyl, d amino acids such as L-d-hydroxylysyl and D-d-methylalanyl, L- $\alpha$ -methylalanyl,  $\beta$  amino acids, and isoquinolyl. D amino acids and non-naturally occurring synthetic amino acids can also be incorporated into the peptides (see, *e.g.*, Roberts *et al.* (1983) *Unusual Amino/Acids in Peptide Synthesis*, 5(6):341-449).

5

10

15

20

25

30

The peptides may also be modified by phosphorylation (see, e.g., W. Bannwarth et al. (1996) Biorganic and Medicinal Chemistry Letters, 6(17):2141-2146), and other methods for making peptide derivatives (see, e.g., Hruby et al. (1990) Biochem. J., 268(2):249-262). Thus, peptide compounds also serve as a basis to prepare peptide mimetics with similar biological activity.

Those of skill in the art recognize that a variety of techniques are available for constructing peptide mimetics with the same or similar desired biological activity as the corresponding peptide compound but with more favorable activity than the peptide with respect to solubility, stability, and susceptibility to hydrolysis and proteolysis (see, e.g., Morgan et al. (1989) An. Rep. Med. Chem., 24:243-252). Methods for preparing peptide mimetics modified at the N-terminal amino group, the C-terminal carboxyl group, and/or changing one or more of the amido linkages in the peptide to a non-amido linkage are known to those of skill in the art.

Amino terminus modifications include alkylating, acetylating, adding a carbobenzoyl group, forming a succinimide group, etc. (see, e.g., Murray et al. (1995) Burger's Medicinal Chemistry and Drug Discovery, 5th ed., Vol. 1, Manfred E. Wolf, ed., John Wiley and Sons, Inc.). C-terminal modifications include mimetics wherein the C-terminal carboxyl group is replaced by an ester, an amide or modifications to form a cyclic peptide.

In addition to N-terminal and C-terminal modifications, the peptide compounds, including peptide mimetics, can advantageously be modified with or covalently coupled to one or more of a variety of hydrophilic polymers. It has been found that when peptide compounds are derivatized with a hydrophilic polymer, their solubility and circulation half-lives may be increased and their

-110-

immunogenicity is masked, with little, if any, diminishment in their binding activity. Suitable nonproteinaceous polymers include, but are not limited to, polyalkylethers as exemplified by polyethylene glycol and polypropylene glycol, polylactic acid, polyglycolic acid, polyoxyalkenes, polyvinylalcohol, polyvinylpyrrolidone, cellulose and cellulose derivatives, dextran and dextran derivatives, etc. Generally, such hydrophilic polymers have an average molecular weight ranging from about 500 to about 100,000 daltons, more preferably from about 2,000 to about 40,000 daltons and, even more preferably, from about 5,000 to about 20,000 daltons. The hydrophilic polymers also can have an average molecular weights of about 5,000 daltons, 10,000 daltons and 20,000 daltons.

10

15

20

25

Methods for derivatizing peptide compounds or for coupling peptides to such polymers have been described (see, e.g., Zallipsky (1995) *Bioconjugate Chem.*, 6:150-165; Monfardini et al. (1995) *Bioconjugate Chem.*, 6:62-69; U.S. Pat. No. 4,640,835; U.S. Pat. No. 4,496,689; U.S. Pat. No. 4,301,144; U.S. Pat. No. 4,670,417; U.S. Pat. No. 4,791,192; U.S. Pat. No. 4,179,337 and WO 95/34326, all of which are incorporated by reference in their entirety herein).

Other methods for making peptide derivatives are described, for example, in Hruby et al. (1990), Biochem J., 268(2):249-262, which is incorporated herein by reference. Thus, the peptide compounds also serve as structural models for non-peptidic compounds with similar biological activity. Those of skill in the art recognize that a variety of techniques are available for constructing compounds with the same or similar desired biological activity as a particular peptide compound but with more favorable activity with respect to solubility, stability, and susceptibility to hydrolysis and proteolysis (see, e.g., Morgan et al. (1989) An. Rep. Med. Chem., 24:243-252, incorporated herein by reference). These techniques include replacing the peptide backbone with a backbone composed of phosphonates, amidates, carbamates, sulfonamides, secondary amines, and N-methylamino acids.

Peptide compounds may exist in a cyclized form with an intramolecular disulfide bond between the thiol groups of the cysteines. Alternatively, an intermolecular disulfide bond between the thiol groups of the cysteines can be

-111-

produced to yield a dimeric (or higher oligomeric) compound. One or more of the cysteine residues may also be substituted with a homocysteine.

# I. CONJUGATES

5

10

15

20

25

30

A conjugate, containing: a) a single chain protease domain (or proteolytically active portion thereof) of an MTSP protein or an MTSP3, MTSP4 or MTSP6 full length zymogen, activated form thereof, or double or single chain protease domain thereof; and b) a targeting agent linked to the MTSP protein directly or via a linker, wherein the agent facilitates: i) affinity isolation or purification of the conjugate; ii) attachment of the conjugate to a surface; iii) detection of the conjugate; or iv) targeted delivery to a selected tissue or cell, is provided herein. The conjugate can be a chemical conjugate or a fusion protein mixture thereof.

The targeting agent is preferably a protein or peptide fragment, such as a tissue specific or tumor specific monoclonal antibody or growth factor or fragment thereof linked either directly or via a linker to an MTSP protein or a protease domain thereof. The targeting agent may also be a protein or peptide fragment that contains a protein binding sequence, a nucleic acid binding sequence, a lipid binding sequence, a polysaccharide binding sequence, or a metal binding sequence, or a linker for attachment to a solid support. In a particular embodiment, the conjugate contains a) the MTSP or portion thereof, as described herein; and b) a targeting agent linked to the MTSP protein directly or via a linker.

Conjugates, such as fusion proteins and chemical conjugates, of the MTSP protein with a protein or peptide fragment (or plurality thereof) that functions, for example, to facilitate affinity isolation or purification of the MTSP protein domain, attachment of the MTSP protein domain to a surface, or detection of the MTSP protein domain are provided. The conjugates can be produced by chemical conjugation, such as via thiol linkages, but are preferably produced by recombinant means as fusion proteins. In the fusion protein, the peptide or fragment thereof is linked to either the N-terminus or C-terminus of the MTSP protein domain. In chemical conjugates the peptide or fragment thereof may be linked anywhere that conjugation can be effected, and there may

-112-

be a plurality of such peptides or fragments linked to a single MTSP protein domain or to a plurality thereof.

The targeting agent is preferably for *in vitro* delivery to a cell or tissue, and includes agents such as cell or tissue-specific antibodies, growth factors and other factors expressed on specific cells; and other cell or tissue specific agents the promote directed delivery of a linked protein.

Most preferably the targeting agent specifically delivers the MTSP protein to selected cells by interaction with a cell surface protein and internalization of conjugate or MTSP protein portion thereof. These conjugate are used in a variety of methods and are particularly suited for use in methods of activation of prodrugs, such as prodrugs that upon cleavage by the particular MTSP protein are cytotoxic. The prodrugs are administered prior to simultaneously with or subsequently to the conjugate. Upon delivery to the targeted cells, the protease activates the prodrug, which then exhibits is therapeutic effect, such as a cytotoxic effect.

# 1. Conjugation

10

15

20

25

30

Conjugates with linked MTSP protein domains can be prepared either by chemical conjugation, recombinant DNA technology, or combinations of recombinant expression and chemical conjugation. The MTSP protein domains and the targeting agent may be linked in any orientation and more than one targeting agents and/or MTSP protein domains may be present in a conjugate.

#### a. Fusion proteins

Fusion proteins are proved herein. A fusion protein contains: a) one or a plurality of domains of an MTSP proteins and b) a targeting agent. The fusion proteins are preferably produced by recombinant expression of nucleic acids that encode the fusion protein.

# b. Chemical conjugation

To effect chemical conjugation herein, the MTSP protein domain is linked via one or more selected linkers or directly to the targeting agent. Chemical conjugation must be used if the targeted agent is other than a peptide or protein, such a nucleic acid or a non-peptide drug. Any means known to those of skill in the art for chemically conjugating selected moieties may be used.

-113-

## 2. Linkers

10

15

20

25

Linkers for two purposes are contemplated herein. The conjugates may include one or more linkers between the MTSP protein portion and the targeting agent. Additionally, linkers are used for facilitating or enhancing immobilization of an MTSP protein or portion thereof on a solid support, such as a microtiter plate, silicon or silicon-coated chip, glass or plastic support, such as for high throughput solid phase screening protocols.

Any linker known to those of skill in the art for preparation of conjugates may be used herein. These linkers are typically used in the preparation of chemical conjugates; peptide linkers may be incorporated into fusion proteins.

Linkers can be any moiety suitable to associate a domain of MTSP protein and a targeting agent. Such linkers and linkages include, but are not limited to, peptidic linkages, amino acid and peptide linkages, typically containing between one and about 60 amino acids, more generally between about 10 and 30 amino acids, chemical linkers, such as heterobifunctional cleavable cross-linkers, including but are not limited to, N-succinimidyl (4-iodoacetyl)-aminobenzoate, sulfosuccinimydil (4-iodoacetyl)-aminobenzoate, 4-succinimidyl-oxycarbonyl-a-(2-pyridyldithio)toluene, sulfosuccinimidyl-6- [a-methyl-a-(pyridyldithiol)toluamido] hexanoate, N-succinimidyl-3-(-2-pyridyldithio) - propionate, succinimidyl 6[3(-(-2-pyridyldithio)-proprionamido] hexanoate, sulfosuccinimidyl 6[3(-(-2-pyridyldithio)-propionamido] hexanoate, 3-(2-pyridyldithio)-propionyl hydrazide, Ellman's reagent, dichlorotriazinic acid, and S-(2-thiopyridyl)-Lcysteine. Other linkers include, but are not limited to peptides and other moieties that reduce stearic hindrance between the domain of MTSP protein and the targeting agent, intracellular enzyme substrates, linkers that increase the flexibility of the conjugate, linkers that increase the solubility of the conjugate, linkers that increase the serum stability of the conjugate, photocleavable linkers and acid cleavable linkers.

-114-

Other exemplary linkers and linkages that are suitable for chemically linked conjugates include, but are not limited to, disulfide bonds, thioether bonds, hindered disulfide bonds, and covalent bonds between free reactive groups, such as amine and thiol groups. These bonds are produced using heterobifunctional reagents to produce reactive thiol groups on one or both of the polypeptides and then reacting the thiol groups on one polypeptide with reactive thiol groups or amine groups to which reactive maleimido groups or thiol groups can be attached on the other. Other linkers include, acid cleavable linkers, such as bismaleimideothoxy propane, acid labile-transferrin conjugates and adipic acid dihydrazide, that would be cleaved in more acidic intracellular compartments; cross linkers that are cleaved upon exposure to UV or visible light and linkers, such as the various domains, such as C<sub>H</sub>1, C<sub>H</sub>2, and C<sub>H</sub>3, from the constant region of human IgG<sub>1</sub> (see, Batra *et al. Molecular Immunol.*, 30:379-386 (1993)). In some embodiments, several linkers may be included in order to take advantage of desired properties of each linker.

10

15

20

25

30

Chemical linkers and peptide linkers may be inserted by covalently coupling the linker to the domain of MTSP protein and the targeting agent. The heterobifunctional agents, described below, may be used to effect such covalent coupling. Peptide linkers may also be linked by expressing DNA encoding the linker and TA, linker and targeted agent, or linker, targeted agent and TA as a fusion protein. Flexible linkers and linkers that increase solubility of the conjugates are contemplated for use, either alone or with other linkers are also contemplated herein.

## a) Acid cleavable, photocleavable and heat sensitive linkers

Acid cleavable linkers, photocleavable and heat sensitive linkers may also be used, particularly where it may be necessary to cleave the domain of MTSP protein to permit it to be more readily accessible to reaction. Acid cleavable linkers include, but are not limited to, bismaleimideothoxy propane; and adipic acid dihydrazide linkers (see, e.g., Fattom et al. (1992) Infection & Immun. 60:584-589) and acid labile transferrin conjugates that contain a sufficient portion of transferrin to permit entry into the intracellular transferrin cycling pathway (see, e.g., Welhöner et al. (1991) J. Biol. Chem. 266:4309-4314).

-115-

Photocleavable linkers are linkers that are cleaved upon exposure to light (see, e.g., Goldmacher et al. (1992) Bioconj. Chem. 3:104-107, which linkers are herein incorporated by reference), thereby releasing the targeted agent upon exposure to light. Photocleavable linkers that are cleaved upon exposure to light are known (see, e.g., Hazum et al. (1981) in Pept., Proc. Eur. Pept. Symp., 16th, Brunfeldt, K (Ed), pp. 105-110, which describes the use of a nitrobenzyl group as a photocleavable protective group for cysteine; Yen et al. (1989) Makromol. Chem 190:69-82, which describes water soluble photocleavable copolymers, including hydroxypropylmethacrylamide copolymer, glycine copolymer, fluorescein copolymer and methylrhodamine copolymer; Goldmacher et al. (1992) Bioconj. Chem. 3:104-107, which describes a cross-linker and reagent that undergoes photolytic degradation upon exposure to near UV light (350 nm); and Senter et al. (1985) Photochem. Photobiol 42:231-237, which describes nitrobenzyloxycarbonyl chloride cross linking reagents that produce photocleavable linkages), thereby releasing the targeted agent upon exposure to light. Such linkers would have particular use in treating dermatological or ophthalmic conditions that can be exposed to light using fiber optics. After administration of the conjugate, the eye or skin or other body part can be exposed to light, resulting in release of the targeted moiety from the conjugate. Such photocleavable linkers are useful in connection with diagnostic protocols in which it is desirable to remove the targeting agent to permit rapid clearance from the body of the animal.

10

15

20

30

#### b) Other linkers for chemical conjugation

Other linkers, include trityl linkers, particularly, derivatized 25 trityl groups to generate a genus of conjugates that provide for release of therapeutic agents at various degrees of acidity or alkalinity. The flexibility thus afforded by the ability to preselect the pH range at which the therapeutic agent will be released allows selection of a linker based on the known physiological differences between tissues in need of delivery of a therapeutic agent (see, e.g., U.S. Patent No. 5,612,474). For example, the acidity of tumor tissues appears to be lower than that of normal tissues.

-116-

## c) Peptide linkers

5

10

15

20

25

30

The linker moieties can be peptides. Peptide linkers can be employed in fusion proteins and also in chemically linked conjugates. The peptide typically has from about 2 to about 60 amino acid residues, for example from about 5 to about 40, or from about 10 to about 30 amino acid residues. The length selected will depend upon factors, such as the use for which the linker is included.

Peptide linkers are advantageous when the targeting agent is proteinaceous. For example, the linker moiety can be a flexible spacer amino acid sequence, such as those known in single-chain antibody research. Examples of such known linker moieties include, but are not limited to, peptides, such as  $(Gly_mSer)_n$  and  $(Ser_mGly)_n$ , in which n is 1 to 6, preferably 1 to 4, more preferably 2 to 4, and m is 1 to 6, preferably 1 to 4, more preferably 2 to 4, enzyme cleavable linkers and others.

Additional linking moieties are described, for example, in Huston *et al.*, *Proc. Natl. Acad. Sci. U.S.A. 85*:5879-5883, 1988; Whitlow, M., *et al.*, *Protein Engineering 6*:989-995, 1993; Newton *et al.*, *Biochemistry 35*:545-553, 1996; A. J. Cumber *et al.*, *Bioconj. Chem. 3*:397-401, 1992; Ladurner *et al.*, *J. Mol. Biol. 273*:330-337, 1997; and U.S. Patent. No. 4,894,443. In some embodiments, several linkers may be included in order to take advantage of desired properties of each linker.

# 3. Targeting agents

Any agent that facilitates detection, immobilization, or purification of the conjugate is contemplated for use herein. For chemical conjugates any moiety that has such properties is contemplated; for fusion proteins, the targeting agent is a protein, peptide or fragment thereof that sufficient to effects the targeting activity. Preferred targeting agents are those that deliver the MTSP protein or portion thereof to selected cells and tissues. Such agents include tumor specific monoclonal antibodies and portions thereof, growth factors, such as FGF, EGF, PDGF, VEGF, cytokines, including chemokines, and other such agents.

# 4. Nucleic acids, plasmids and cells

-117-

Isolated nucleic acid fragments encoding fusion proteins are provided. The nucleic acid fragment that encodes the fusion protein includes: a) nucleic acid encoding a protease domain of an MTSP protein encoded by a nucleic acid that hybridizes to a nucleic acid having the nucleotide sequence set forth in the SEQ. ID NO:1; and b) nucleic acid encoding a protein, peptide or effective fragment thereof that facilitates: i) affinity isolation or purification of the fusion protein; ii) attachment of the fusion protein to a surface; or iii) detection of the fusion protein. Preferably, the nucleic acid is DNA..

Plasmids for replication and vectors for expression that contain the above nucleic acid fragments are also provided. Cells containing the plasmids and vectors are also provided. The cells can be any suitable host including, but are not limited to, bacterial cells, yeast cells, fungal cells, plant cells, insect cell and animal cells. The nucleic acids, plasmids, and cells containing the plasmids can be prepared according to methods known in the art including any described herein.

10

15

20

25

30

Also provided are methods for producing the above fusion proteins. An exemplary method includes the steps of growing, i.e. culturing the cells so that the proliferate, cells containing a plasmid encoding the fusion protein under conditions whereby the fusion protein is expressed by the cell, and recovering the expressed fusion protein. Methods for expressing and recovering recombinant proteins are well known in the art (*See generally, Current Protocols in Molecular Biology* (1998) § 16, John Wiley & Sons, Inc.) and such methods can be used for expressing and recovering the expressed fusion proteins. Preferably, the recombinant expression and recovery methods disclosed in Section B can be used.

The recovered fusion proteins can be isolated or purified by methods known in the art such as centrifugation, filtration, chromatograph, electrophoresis, immunoprecipitation, etc., or by a combination thereof (*See generally, Current Protocols in Molecular Biology* (1998) § 10, John Wiley & Sons, Inc.). Preferably, the recovered fusion protein is isolated or purified through affinity binding between the protein or peptide fragment of the fusion protein and an affinity binding moiety. As discussed in the above sections

-118-

regarding the construction of the fusion proteins, any affinity binding pairs can be constructed and used in the isolation or purification of the fusion proteins. For example, the affinity binding pairs can be protein binding sequences/protein, DNA binding sequences/DNA sequences, RNA binding sequences/RNA sequences, lipid binding sequences/lipid, polysaccharide binding sequences/polysaccharide, or metal binding sequences/metal.

# 5. Immobilization and supports or substrates therefor

10

15

20

25

30

In certain embodiments, where the targeting agents are designed for linkage to surfaces, the MTSP protein can be attached by linkage such as ionic or covalent, non-covalent or other chemical interaction, to a surface of a support or matrix material. Immobilization may be effected directly or via a linker. The MTSP protein may be immobilized on any suitable support, including, but are not limited to, silicon chips, and other supports described herein and known to those of skill in the art. A plurality of MTSP protein or protease domains thereof may be attached to a support, such as an array (*i.e.*, a pattern of two or more) of conjugates on the surface of a silicon chip or other chip for use in high throughput protocols and formats.

It is also noted that the domains of the MTSP protein can be linked directly to the surface or via a linker without a targeting agent linked thereto. Hence chips containing arrays of the domains of the MTSP protein.

The matrix material or solid supports contemplated herein are generally any of the insoluble materials known to those of skill in the art to immobilize ligands and other molecules, and are those that used in many chemical syntheses and separations. Such supports are used, for example, in affinity chromatography, in the immobilization of biologically active materials, and during chemical syntheses of biomolecules, including proteins, amino acids and other organic molecules and polymers. The preparation of and use of supports is well known to those of skill in this art; there are many such materials and preparations thereof known. For example, naturally-occurring support materials, such as agarose and cellulose, may be isolated from their respective sources, and processed according to known protocols, and synthetic materials may be prepared in accord with known protocols.

-119-

The supports are typically insoluble materials that are solid, porous, deformable, or hard, and have any required structure and geometry, including, but not limited to: beads, pellets, disks, capillaries, hollow fibers, needles, solid fibers, random shapes, thin films and membranes. Thus, the item may be fabricated from the matrix material or combined with it, such as by coating all or part of the surface or impregnating particles.

Typically, when the matrix is particulate, the particles are at least about  $10\text{-}2000~\mu\text{M}$ , but may be smaller or larger, depending upon the selected application. Selection of the matrices will be governed, at least in part, by their physical and chemical properties, such as solubility, functional groups, mechanical stability, surface area swelling propensity, hydrophobic or hydrophilic properties and intended use.

10

15

20

25

30

If necessary, the support matrix material can be treated to contain an appropriate reactive moiety. In some cases, the support matrix material already containing the reactive moiety may be obtained commercially. The support matrix material containing the reactive moiety may thereby serve as the matrix support upon which molecules are linked. Materials containing reactive surface moieties such as amino silane linkages, hydroxyl linkages or carboxysilane linkages may be produced by well established surface chemistry techniques involving silanization reactions, or the like. Examples of these materials are those having surface silicon oxide moieties, covalently linked to gamma-aminopropylsilane, and other organic moieties; N-[3-(triethyoxysilyl)propyl]phthelamic acid; and bis-(2-hydroxyethyl)aminopropyltriethoxysilane. Exemplary of readily available materials containing amino group reactive functionalities, include, but are not limited to, para-aminophenyltriethyoxysilane. Also derivatized polystyrenes and other such polymers are well known and readily available to those of skill in this art (e.g., the Tentagel® Resins are available with a multitude of functional groups, and are sold by Rapp Polymere, Tubingen, Germany; see, U.S. Patent No. 4,908,405 and U.S. Patent No. 5,292,814; see, also Butz et al., Peptide Res., 7:20-23 (1994); and Kleine et al., Immunobiol., 190:53-66 (1994)).

-120-

These matrix materials include any material that can act as a support matrix for attachment of the molecules of interest. Such materials are known to those of skill in this art, and include those that are used as a support matrix. These materials include, but are not limited to, inorganics, natural polymers, and synthetic polymers; including, but are not limited to: cellulose, cellulose derivatives, acrylic resins, glass, silica gels, polystyrene, gelatin, polyvinyl pyrrolidone, co-polymers of vinyl and acrylamide, polystyrene cross-linked with divinylbenzene and others (see, Merrifield, Biochemistry, 3:1385-1390 (1964)), polyacrylamides, latex gels, polystyrene, dextran, polyacrylamides, rubber, silicon, plastics, nitrocellulose, celluloses, natural sponges. Of particular interest herein, are highly porous glasses (see, e.g., U.S. Patent No. 4,244,721) and others prepared by mixing a borosilicate, alcohol and water.

10

Synthetic supports include, but are not limited to: acrylamides, dextranderivatives and dextran co-polymers, agarose-polyacrylamide blends, other polymers and co-polymers with various functional groups, methacrylate 15 derivatives and co-polymers, polystyrene and polystyrene copolymers (see, e.g., Merrifield, Biochemistry, 3:1385-1390 (1964); Berg et al., in Innovation Perspect. Solid Phase Synth. Collect. Pap., Int. Symp., 1st, Epton, Roger (Ed), pp. 453-459 (1990); Berg et al., Pept., Proc. Eur. Pept. Symp., 20th, Jung, G. 20 et al. (Eds), pp. 196-198 (1989); Berg et al., J. Am. Chem. Soc., 111:8024-8026 (1989); Kent et al., Isr. J. Chem., 17:243-247 (1979); Kent et al., J. Org. Chem., 43:2845-2852 (1978); Mitchell et al., Tetrahedron Lett., 42:3795-3798 (1976); U.S. Patent No. 4,507,230; U.S. Patent No. 4,006,117; and U.S. Patent No. 5,389,449). Such materials include those made from 25 polymers and co-polymers such as polyvinylalcohols, acrylates and acrylic acids such as polyethylene-co-acrylic acid, polyethylene-co-methacrylic acid, polyethylene-co-ethylacrylate, polyethylene-co-methyl acrylate, polypropylene-co-acrylic acid, polypropylene-co-methyl-acrylic acid, polypropylene-co-ethylacrylate, polypropylene-co-methyl acrylate, polyethylene-co-vinyl acetate, poly-30 propylene-co-vinyl acetate, and those containing acid anhydride groups such as polyethylene-co-maleic anhydride and polypropylene-co-maleic anhydride.

-121-

Liposomes have also been used as solid supports for affinity purifications (Powell et al. *Biotechnol. Bioeng.*, <u>33</u>:173 (1989)).

Numerous methods have been developed for the immobilization of proteins and other biomolecules onto solid or liquid supports (see, e.g.,

5 Mosbach, Methods in Enzymology, 44 (1976); Weetall, Immobilized Enzymes, Antigens, Antibodies, and Peptides, (1975); Kennedy et al., Solid Phase Biochemistry, Analytical and Synthetic Aspects, Scouten, ed., pp. 253-391 (1983); see, generally, Affinity Techniques. Enzyme Purification: Part B. Methods in Enzymology, Vol. 34, ed. W. B. Jakoby, M. Wilchek, Acad. Press, N.Y. (1974); and Immobilized Biochemicals and Affinity Chromatography, Advances in Experimental Medicine and Biology, vol. 42, ed. R. Dunlap, Plenum Press, N.Y. (1974)).

Among the most commonly used methods are absorption and adsorption or covalent binding to the support, either directly or via a linker, such as the numerous disulfide linkages, thioether bonds, hindered disulfide bonds, and covalent bonds between free reactive groups, such as amine and thiol groups, known to those of skill in art (see, e.g., the PIERCE CATALOG, ImmunoTechnology Catalog & Handbook, 1992-1993, which describes the preparation of and use of such reagents and provides a commercial source for such reagents; Wong, Chemistry of Protein Conjugation and Cross Linking, CRC Press (1993); see also DeWitt et al., Proc. Natl. Acad. Sci. U.S.A., 90:6909 (1993); Zuckermann et al., J. Am. Chem. Soc., 114:10646 (1992); Kurth et al., J. Am. Chem. Soc., 116:2661 (1994); Ellman et al., Proc. Natl. Acad. Sci. U.S.A., 91:4708 (1994); Sucholeiki, Tetrahedron Lttrs., 35:7307 (1994); Su-Sun Wang, J. Org. Chem., 41:3258 (1976); Padwa et al., J. Org. Chem., 41:3550 (1971); and Vedejs et al., J. Org. Chem., 49:575 (1984), which describe photosensitive linkers).

15

20

25

30

To effect immobilization, a composition containing the protein or other biomolecule is contacted with a support material such as alumina, carbon, an ion-exchange resin, cellulose, glass or a ceramic. Fluorocarbon polymers have been used as supports to which biomolecules have been attached by adsorption

-122-

(see, U.S. Patent No. 3,843,443; Published International PCT Application WO/86 03840).

# J. Prognosis and diagnosis

5

10

15

20

25

30

MTSP protein proteins, domains, analogs, and derivatives thereof, and encoding nucleic acids (and sequences complementary thereto), and anti-MTSP protein antibodies, can be used in diagnostics. Such molecules can be used in assays, such as immunoassays, to detect, prognose, diagnose, or monitor various conditions, diseases, and disorders affecting MTSP protein expression, or monitor the treatment thereof. For purposes herein, the presence of MTSPs in body fluids or tumor tissues are of particular interest.

In particular, such an immunoassay is carried out by a method including contacting a sample derived from a patient with an anti-MTSP protein antibody under conditions such that specific binding can occur, and detecting or measuring the amount of any specific binding by the antibody. In a specific aspect, such binding of antibody, in tissue sections, can be used to detect aberrant MTSP protein localization or aberrant (e.g., low or absent) levels of MTSP protein. In a specific embodiment, antibody to MTSP protein can be used to assay in a patient tissue or serum sample for the presence of MTSP protein where an aberrant level of MTSP protein is an indication of a diseased condition.

The immunoassays which can be used include but are not limited to competitive and non-competitive assay systems using techniques such as western blots, radioimmunoassays, ELISA (enzyme linked immunosorbent assay), "sandwich" immunoassays, immunoprecipitation assays, precipitin reactions, gel diffusion precipitin reactions, immunodiffusion assays, agglutination assays, complement-fixation assays, immunoradiometric assays, fluorescent immunoassays, protein A immunoassays, to name but a few.

MTSP protein genes and related nucleic acid sequences and subsequences, including complementary sequences, can also be used in hybridization assays. MTSP protein nucleic acid sequences, or subsequences thereof containing about at least 8 nucleotides, preferably 14 or 16 or 30 or more continugous nucleotides, can be used as hybridization probes. Hybridization assays can be used to detect, prognose, diagnose, or monitor

-123-

conditions, disorders, or disease states associated with aberrant changes in MTSP protein expression and/or activity as described herein. In particular, such a hybridization assay is carried out by a method by contacting a sample containing nucleic acid with a nucleic acid probe capable of hybridizing to MTSP protein encoding DNA or RNA, under conditions such that hybridization can occur, and detecting or measuring any resulting hybridization.

5

10

15

20

25

30

In a specific embodiment, a method of diagnosing a disease or disorder characterized by detecting an aberrant level of an MTSP protein in a subject is provided herein by measuring the level of the DNA, RNA, protein or functional activity of the epithelial MTSP protein at least partially encoded by a nucleic acid that hybridizes to a nucleic acid having the nucleotide sequence set forth in the SEQ. ID NO:1 in a sample derived from the subject, wherein an increase or decrease in the level of the DNA, RNA, protein or functional activity of the MTSP protein, relative to the level of the DNA, RNA, protein or functional activity found in an analogous sample not having the disease or disorder indicates the presence of the disease or disorder in the subject.

Kits for diagnostic use are also provided, that contain in one or more containers an anti-MTSP protein antibody, particularly anti-MTSP3 or anti = MTSP4, and, optionally, a labeled binding partner to the antibody. Alternatively, the anti-MTSP protein antibody can be labeled (with a detectable marker, e.g., a chemiluminescent, enzymatic, fluorescent, or radioactive moiety). A kit is also provided that includes in one or more containers a nucleic acid probe capable of hybridizing to MTSP protein-encoding RNA. In a specific embodiment, a kit can comprise in one or more containers a pair of primers (e.g., each in the size range of 6-30 nucleotides) that are capable of priming amplification [e.g., by polymerase chain reaction (see e.g., Innis et al., 1990, PCR Protocols, Academic Press, Inc., San Diego, CA), ligase chain reaction (see EP 320,308) use of  $\Omega\beta$  replicase, cyclic probe reaction, or other methods known in the art under appropriate reaction conditions of at least a portion of an MTSP protein-encoding nucleic acid. A kit can optionally further comprise in a container a predetermined amount of a purified MTSP protein or nucleic acid, e.g., for use as a standard or control.

-124-

# K. PHARMACEUTICAL COMPOSITIONS AND MODES OF ADMINISTRATION

## 1. Components of the compositions

5

10

15

20

25

30

Pharmaceutical compositions containing the identified compounds that modulate the activity of an MTSP protein are provided herein. Also provided are combinations of a compound that modulates the activity of an MTSP protein and another treatment or compound for treatment of a neoplastic disorder, such as a chemotherapeutic compound.

The MTSP protein modulator and the anti-tumor agent can be packaged as separate compositions for administration together or sequentially or intermittently. Alternatively, they can provided as a single composition for administration or as two compositions for administration as a single composition. The combinations can be packaged as kits.

## a. MTSP protein inhibitors

Any MTSP protein inhibitors, including those described herein when used alone or in combination with other compounds, that can alleviate, reduce, ameliorate, prevent, or place or maintain in a state of remission of clinical symptoms or diagnostic markers associated with neoplastic diseases, including undesired and/or uncontrolled angiogenesis, can be used in the present combinations.

In one embodiment, the MTSP protein inhibitor is an antibody or fragment thereof that specifically reacts with an MTSP protein or the protease domain thereof, an inhibitor of the MTSP protein production, an inhibitor of the epithelial MTSP protein membrane-localization, or any inhibitor of the expression of or, especially, the activity of an MTSP protein.

## b. Anti-angiogenic agents and anti-tumor agents

Any anti-angiogenic agents and anti-tumor agents, including those described herein, when used alone or in combination with other compounds, that can alleviate, reduce, ameliorate, prevent, or place or maintain in a state of remission of clinical symptoms or diagnostic markers associated with undesired and/or uncontrolled angiogenesis and/or tumor growth and metastasis, particularly solid neoplasms, vascular malformations and cardiovascular disorders, chronic inflammatory diseases and aberrant wound repairs, circulatory

-125-

disorders, crest syndromes, dermatological disorders, or ocular disorders, can be used in the combinations. Also contemplated are anti-tumor agents for use in combination with an inhibitor of an MTSP protein.

## c. Anti-tumor agents and anti-angiogenic agents

The compounds identified by the methods provided herein or provided herein can be used in combination with anti-tumor agents and/or anti-angiogenesis agents.

## 2. Formulations and route of administration

5

10

15

20

25

30

The compounds herein and agents are preferably formulated as pharmaceutical compositions, preferably for single dosage administration. The concentrations of the compounds in the formulations are effective for delivery of an amount, upon administration, that is effective for the intended treatment. Typically, the compositions are formulated for single dosage administration. To formulate a composition, the weight fraction of a compound or mixture thereof is dissolved, suspended, dispersed or otherwise mixed in a selected vehicle at an effective concentration such that the treated condition is relieved or ameliorated. Pharmaceutical carriers or vehicles suitable for administration of the compounds provided herein include any such carriers known to those skilled in the art to be suitable for the particular mode of administration.

In addition, the compounds may be formulated as the sole pharmaceutically active ingredient in the composition or may be combined with other active ingredients. Liposomal suspensions, including tissue-targeted liposomes, may also be suitable as pharmaceutically acceptable carriers. These may be prepared according to methods known to those skilled in the art. For example, liposome formulations may be prepared as described in U.S. Patent No. 4,522,811.

The active compound is included in the pharmaceutically acceptable carrier in an amount sufficient to exert a therapeutically useful effect in the absence of undesirable side effects on the patient treated. The therapeutically effective concentration may be determined empirically by testing the compounds in known in vitro and in vivo systems, such as the assays provided herein.

-126-

The concentration of active compound in the drug composition will depend on absorption, inactivation and excretion rates of the active compound, the physicochemical characteristics of the compound, the dosage schedule, and amount administered as well as other factors known to those of skill in the art.

5

10

15

20

25

Typically a therapeutically effective dosage is contemplated. The amounts administered may be on the order of 0.001 to 1 mg/ml, preferably about 0.005-0.05 mg/ml, more preferably about 0.01 mg/ml, of blood volume. Pharmaceutical dosage unit forms are prepared to provide from about 1 mg to about 1000 mg and preferably from about 10 to about 500 mg, more preferably about 25-75 mg of the essential active ingredient or a combination of essential ingredients per dosage unit form. The precise dosage can be empirically determined.

The active ingredient may be administered at once, or may be divided into a number of smaller doses to be administered at intervals of time. It is understood that the precise dosage and duration of treatment is a function of the disease being treated and may be determined empirically using known testing protocols or by extrapolation from in vivo or in vitro test data. It is to be noted that concentrations and dosage values may also vary with the severity of the condition to be alleviated. It is to be further understood that for any particular subject, specific dosage regimens should be adjusted over time according to the individual need and the professional judgment of the person administering or supervising the administration of the compositions, and that the concentration ranges set forth herein are exemplary only and are not intended to limit the scope or use of the claimed compositions and combinations containing them.

Preferred pharmaceutically acceptable derivatives include acids, salts, esters, hydrates, solvates and prodrug forms. The derivative is typically selected such that its pharmacokinetic properties are superior to the corresponding neutral compound.

Thus, effective concentrations or amounts of one or more of the compounds provided herein or pharmaceutically acceptable derivatives thereof are mixed with a suitable pharmaceutical carrier or vehicle for systemic, topical or local administration to form pharmaceutical compositions. Compounds are included in an amount effective for ameliorating or treating the disorder for which treatment is contemplated. The concentration of active compound in the composition will depend on absorption, inactivation, excretion rates of the active compound, the dosage schedule, amount administered, particular formulation as well as other factors known to those of skill in the art.

Solutions or suspensions used for parenteral, intradermal, subcutaneous, or topical application can include any of the following components: a sterile diluent, such as water for injection, saline solution, fixed oil, polyethylene glycol, glycerine, propylene glycol or other synthetic solvent; antimicrobial agents, such as benzyl alcohol and methyl parabens; antioxidants, such as ascorbic acid and sodium bisulfite; chelating agents, such as ethylenediaminetetraacetic acid (EDTA); buffers, such as acetates, citrates and phosphates; and agents for the adjustment of tonicity such as sodium chloride or dextrose. Parenteral preparations can be enclosed in ampules, disposable syringes or single or multiple dose vials made of glass, plastic or other suitable material.

10

15

20

25

30

In instances in which the compounds exhibit insufficient solubility, methods for solubilizing compounds may be used. Such methods are known to those of skill in this art, and include, but are not limited to, using cosolvents, such as dimethylsulfoxide (DMSO), using surfactants, such as Tween<sup>®</sup>, or dissolution in aqueous sodium bicarbonate. Derivatives of the compounds, such as prodrugs of the compounds may also be used in formulating effective pharmaceutical compositions. For ophthalmic indications, the compositions are formulated in an ophthalmically acceptable carrier. For the ophthalmic uses herein, local administration, either by topical administration or by injection is preferred. Time release formulations are also desirable. Typically, the compositions are formulated for single dosage administration, so that a single dose administers an effective amount.

-128-

Upon mixing or addition of the compound with the vehicle, the resulting mixture may be a solution, suspension, emulsion or other composition. The form of the resulting mixture depends upon a number of factors, including the intended mode of administration and the solubility of the compound in the selected carrier or vehicle. If necessary, pharmaceutically acceptable salts or other derivatives of the compounds are prepared.

The compound is included in the pharmaceutically acceptable carrier in an amount sufficient to exert a therapeutically useful effect in the absence of undesirable side effects on the patient treated. It is understood that number and degree of side effects depends upon the condition for which the compounds are administered. For example, certain toxic and undesirable side effects are tolerated when treating life-threatening illnesses that would not be tolerated when treating disorders of lesser consequence.

10

15

20

25

30

The compounds can also be mixed with other active materials, that do not impair the desired action, or with materials that supplement the desired action known to those of skill in the art. The formulations of the compounds and agents for use herein include those suitable for oral, rectal, topical, inhalational, buccal (e.g., sublingual), parenteral (e.g., subcutaneous, intramuscular, intradermal, or intravenous), transdermal administration or any route. The most suitable route in any given case will depend on the nature and severity of the condition being treated and on the nature of the particular active compound which is being used. The formulations are provided for administration to humans and animals in unit dosage forms, such as tablets, capsules, pills, powders, granules, sterile parenteral solutions or suspensions, and oral solutions or suspensions, and oil-water emulsions containing suitable quantities of the compounds or pharmaceutically acceptable derivatives thereof. The pharmaceutically therapeutically active compounds and derivatives thereof are typically formulated and administered in unit-dosage forms or multiple-dosage forms. Unit-dose forms as used herein refers to physically discrete units suitable for human and animal subjects and packaged individually as is known in the art. Each unit-dose contains a predetermined quantity of the therapeutically active compound sufficient to produce the desired therapeutic effect, in association

-129-

with the required pharmaceutical carrier, vehicle or diluent. Examples of unit-dose forms include ampoules and syringes and individually packaged tablets or capsules. Unit-dose forms may be administered in fractions or multiples thereof. A multiple-dose form is a plurality of identical unit-dosage forms packaged in a single container to be administered in segregated unit-dose form. Examples of multiple-dose forms include vials, bottles of tablets or capsules or bottles of pints or gallons. Hence, multiple dose form is a multiple of unit-doses which are not segregated in packaging.

The composition can contain along with the active ingredient: a diluent 10 such as lactose, sucrose, dicalcium phosphate, or carboxymethylcellulose; a lubricant, such as magnesium stearate, calcium stearate and talc; and a binder such as starch, natural gums, such as gum acaciagelatin, glucose, molasses, polvinylpyrrolidine, celluloses and derivatives thereof, povidone, crospovidones and other such binders known to those of skill in the art. Liquid 15 pharmaceutically administrable compositions can, for example, be prepared by dissolving, dispersing, or otherwise mixing an active compound as defined above and optional pharmaceutical adjuvants in a carrier, such as, for example, water, saline, aqueous dextrose, glycerol, glycols, ethanol, and the like, to thereby form a solution or suspension. If desired, the pharmaceutical composition to be 20 administered may also contain minor amounts of nontoxic auxiliary substances such as wetting agents, emulsifying agents, or solubilizing agents, pH buffering agents and the like, for example, acetate, sodium citrate, cyclodextrine derivatives, sorbitan monolaurate, triethanolamine sodium acetate, triethanolamine oleate, and other such agents. Methods of preparing such 25 dosage forms are known, or will be apparent, to those skilled in this art (see, e.g., Remington's Pharmaceutical Sciences, Mack Publishing Company, Easton, Pa., 15th Edition, 1975). The composition or formulation to be administered will contain a quantity of the active compound in an amount sufficient to alleviate the symptoms of the treated subject.

Dosage forms or compositions containing active ingredient in the range of 0.005% to 100% with the balance made up from non-toxic carrier may be prepared. For oral administration, the pharmaceutical compositions may take the

30

-130-

form of, for example, tablets or capsules prepared by conventional means with pharmaceutically acceptable excipients such as binding agents (e.g., pregelatinized maize starch, polyvinyl pyrrolidone or hydroxypropyl methylcellulose); fillers (e.g., lactose, microcrystalline cellulose or calcium hydrogen phosphate); lubricants (e.g., magnesium stearate, talc or silica); disintegrants (e.g., potato starch or sodium starch glycolate); or wetting agents (e.g., sodium lauryl sulphate). The tablets may be coated by methods well-known in the art.

The pharmaceutical preparation may also be in liquid form, for example, solutions, syrups or suspensions, or may be presented as a drug product for reconstitution with water or other suitable vehicle before use. Such liquid preparations may be prepared by conventional means with pharmaceutically acceptable additives such as suspending agents (e.g., sorbitol syrup, cellulose derivatives or hydrogenated edible fats); emulsifying agents (e.g., lecithin or acacia); non-aqueous vehicles (e.g., almond oil, oily esters, or fractionated vegetable oils); and preservatives (e.g., methyl or propyl-p-hydroxybenzoates or sorbic acid).

10

15

20

25

30

Formulations suitable for rectal administration are preferably presented as unit dose suppositories. These may be prepared by admixing the active compound with one or more conventional solid carriers, for example, cocoa butter, and then shaping the resulting mixture.

Formulations suitable for topical application to the skin or to the eye preferably take the form of an ointment, cream, lotion, paste, gel, spray, aerosol and oil. Carriers which may be used include vaseline, lanoline, polyethylene glycols, alcohols, and combinations of two or more thereof. The topical formulations may further advantageously contain 0.05 to 15 percent by weight of thickeners selected from among hydroxypropyl methyl cellulose, methyl cellulose, polyvinylpyrrolidone, polyvinyl alcohol, poly (alkylene glycols), poly/hydroxyalkyl, (meth)acrylates or poly(meth)acrylamides. A topical formulation is often applied by instillation or as an ointment into the conjunctival sac. It can also be used for irrigation or lubrication of the eye, facial sinuses, and external auditory meatus. It may also be injected into the anterior eye

chamber and other places. The topical formulations in the liquid state may be also present in a hydrophilic three-dimensional polymer matrix in the form of a strip, contact lens, and the like from which the active components are released.

5

10

15

20

25

30

For administration by inhalation, the compounds for use herein can be delivered in the form of an aerosol spray presentation from pressurized packs or a nebulizer, with the use of a suitable propellant, *e.g.*, dichlorodifluoromethane, trichlorofluoromethane, dichlorotetrafluoroethane, carbon dioxide or other suitable gas. In the case of a pressurized aerosol, the dosage unit may be determined by providing a valve to deliver a metered amount. Capsules and cartridges of, *e.g.*, gelatin, for use in an inhaler or insufflator may be formulated containing a powder mix of the compound and a suitable powder base such as lactose or starch.

Formulations suitable for buccal (sublingual) administration include, for example, lozenges containing the active compound in a flavored base, usually sucrose and acacia or tragacanth; and pastilles containing the compound in an inert base such as gelatin and glycerin or sucrose and acacia.

The compounds may be formulated for parenteral administration by injection, e.g., by bolus injection or continuous infusion. Formulations for injection may be presented in unit dosage form, e.g., in ampules or in multi-dose containers, with an added preservative. The compositions may be suspensions, solutions or emulsions in oily or aqueous vehicles, and may contain formulatory agents such as suspending, stabilizing and/or dispersing agents. Alternatively, the active ingredient may be in powder form for reconstitution with a suitable vehicle, e.g., sterile pyrogen-free water or other solvents, before use.

Formulations suitable for transdermal administration may be presented as discrete patches adapted to remain in intimate contact with the epidermis of the recipient for a prolonged period of time. Such patches suitably contain the active compound as an optionally buffered aqueous solution of, for example, 0.1 to 0.2 M concentration with respect to the active compound. Formulations suitable for transdermal administration may also be delivered by iontophoresis (see, e.g., Pharmaceutical Research 3 (6), 318 (1986)) and typically take the form of an optionally buffered aqueous solution of the active compound.

The pharmaceutical compositions may also be administered by controlled release means and/or delivery devices (see, *e.g.*, in U.S. Patent Nos. 3,536,809; 3,598,123; 3,630,200; 3,845,770; 3,847,770; 3,916,899; 4,008,719; 4,687,610; 4,769,027; 5,059,595; 5,073,543; 5,120,548; 5,354,566; 5,591,767; 5,639,476; 5,674,533 and 5,733,566).

5

10

15

20

25

30

Desirable blood levels may be maintained by a continuous infusion of the active agent as ascertained by plasma levels. It should be noted that the attending physician would know how to and when to terminate, interrupt or adjust therapy to lower dosage due to toxicity, or bone marrow, liver or kidney dysfunctions. Conversely, the attending physician would also know how to and when to adjust treatment to higher levels if the clinical response is not adequate (precluding toxic side effects).

The efficacy and/or toxicity of the MTSP protein inhibitor(s), alone or in combination with other agents can also be assessed by the methods known in the art (See generally, O'Reilly, *Investigational New Drugs*, <u>15</u>:5-13 (1997)).

The active compounds or pharmaceutically acceptable derivatives may be prepared with carriers that protect the compound against rapid elimination from the body, such as time release formulations or coatings.

Kits containing the compositions and/or the combinations with instructions for administration thereof are provided. The kit may further include a needle or syringe, preferably packaged in sterile form, for injecting the complex, and/or a packaged alcohol pad. Instructions are optionally included for administration of the active agent by a clinician or by the patient.

Finally, the compounds or MTSP proteins or protease domains thereof or compositions containing any of the preceding agents may be packaged as articles of manufacture containing packaging material, a compound or suitable derivative thereof provided herein, which is effective for treatment of a diseases or disorders contemplated herein, within the packaging material, and a label that indicates that the compound or a suitable derivative thereof is for treating the diseases or disorders contemplated herein. The label can optionally include the disorders for which the therapy is warranted.

-133-

## L. METHODS OF TREATMENT

5

15

20

25

30

The compounds identified by the methods herein are used for treating or preventing neoplastic diseases in an animal, particularly a mammal, including a human, is provided herein. In one embodiment, the method includes administering to a mammal an effective amount of an inhibitor of an MTSP protein, whereby the disease or disorder is treated or prevented. In a preferred embodiment, the MTSP protein inhibitor used in the treatment or prevention is administered with a pharmaceutically acceptable carrier or excipient. The mammal treated can be a human.

10 The inhibitors provided herein are those identified by the screening assays. In addition, antibodies and antisense nucleic acids are contemplated.

The treatment or prevention method can further include administering an anti-angiogenic treatment or agent or anti-tumor agent simultaneously with, prior to or subsequent to the MTSP protein inhibitor, which can be any compound identified that inhibits the activity of an MTSP protein, and includes an antibody or a fragment or derivative thereof containing the binding region thereof against the MTSP protein, an antisense nucleic acid encoding the MTSP protein, and a nucleic acid containing at least a portion of a gene encoding the MTSP protein into which a heterologous nucleotide sequence has been inserted such that the heterologous sequence inactivates the biological activity of at least a portion of the gene encoding the MTSP protein, in which the portion of the gene encoding the MTSP protein flanks the heterologous sequence so as to promote homologous recombination with a genomic gene encoding the MTSP protein.

## 1. Antisense treatment

In a specific embodiment, as described hereinabove, MTSP protein function is reduced or inhibited by MTSP protein antisense nucleic acids, to treat or prevent neoplastic disease. The therapeutic or prophylactic use of nucleic acids of at least six nucleotides that are antisense to a gene or cDNA encoding MTSP protein or a portion thereof. An MTSP protein "antisense" nucleic acid as used herein refers to a nucleic acid capable of hybridizing to a portion of an MTSP protein RNA (preferably mRNA) by virtue of some sequence complementarily. The antisense nucleic acid may be complementary to a coding

and/or noncoding region of an MTSP protein mRNA. Such antisense nucleic acids have utility as therapeutics that reduce or inhibit MTSP protein function, and can be used in the treatment or prevention of disorders as described *supra*.

5

10

15

20

25

30

The MTSP protein antisense nucleic acids are of at least six nucleotides and are preferably oligonucleotides (ranging from 6 to about 150 nucleotides, or more preferably 6 to 50 nucleotides). In specific aspects, the oligonucleotide is at least 10 nucleotides, at least 15 nucleotides, at least 100 nucleotides, or at least 125 nucleotides. The oligonucleotides can be DNA or RNA or chimeric mixtures or derivatives or modified versions thereof, single-stranded or doublestranded. The oligonucleotide can be modified at the base moiety, sugar moiety, or phosphate backbone. The oligonucleotide may include other appending groups such as peptides, or agents facilitating transport across the cell membrane (see, e.g., Letsinger et al., Proc. Natl. Acad. Sci. U.S.A. 86:6553-6556 (1989); Lemaitre et al., *Proc. Natl. Acad. Sci. U.S.A.* <u>84</u>:648-652 (1987); PCT Publication No. WO 88/09810, published December 15, 1988) or bloodbrain barrier (see, e.g., PCT Publication No. WO 89/10134, published April 25, 1988), hybridization-triggered cleavage agents (see, e.g., Krol et al., BioTechniques 6:958-976 (1988)) or intercalating agents (see, e.g., Zon, Pharm. Res. 5:539-549 (1988)).

The MTSP protein antisense nucleic acid is preferably an oligonucleotide, more preferably of single-stranded DNA. In a preferred aspect, the oligonucleotide includes a sequence antisense to a portion of human MTSP protein. The oligonucleotide may be modified at any position on its structure with substituents generally known in the art.

The MTSP protein antisense oligonucleotide may comprise at least one modified base moiety which is selected from the group including, but not limited to 5-fluorouracil, 5-bromouracil, 5-chlorouracil, 5-iodouracil, hypoxanthine, xanthine, 4-acetylcytosine, 5-(carboxyhydroxylmethyl) uracil, 5-carboxymethylaminomethyl-2-thiouridine, 5-carboxymethylaminomethyluracil, dihydrouracil, beta-D-galactosylqueosine, inosine, N6-isopentenyladenine, 1-methylguanine, 1-methylguanine, 2,2-dimethylguanine, 2-methyladenine, 2-methylguanine, 3-methylcytosine, 5-methylcytosine, N6-adenine,

7-methylguanine, 5-methylaminomethyluracil, 5-methoxyaminomethyl2-thiouracil, beta-D-mannosylqueosine, 5'-methoxycarboxymethyluracil,
5-methoxyuracil, 2-methylthio-N6-isopentenyladenine, uracil-5-oxyacetic acid (v),
wybutoxosine, pseudouracil, queosine, 2-thiocytosine, 5-methyl-2-thiouracil,
2-thiouracil, 4-thiouracil, 5-methyluracil, uracil-5-oxyacetic acid methylester,
uracil-5-oxyacetic acid (v), 5-methyl-2-thiouracil, 3-(3-amino-3-N-2carboxypropyl) uracil, (acp3)w, and 2,6-diaminopurine.

In another embodiment, the oligonucleotide includes at least one modified sugar moiety selected from the group including but not limited to arabinose,

2-fluoroarabinose, xylulose, and hexose. The oligonucleotide can include at least one modified phosphate backbone selected from a phosphorothioate, a phosphorodithioate, a phosphoramidothioate, a phosphoramidate, a phosphordiamidate, a methylphosphonate, an alkyl phosphotriester, and a formacetal or analog thereof.

The oligonucleotide can be an  $\alpha$ -anomeric oligonucleotide. An  $\alpha$ -anomeric oligonucleotide forms specific double-stranded hybrids with complementary RNA in which the strands run parallel to each other (Gautier et al., *Nucl. Acids Res.* 15:6625-6641 (1987)).

The oligonucleotide may be conjugated to another molecule, *e.g.*, a peptide, hybridization triggered cross-linking agent, transport agent and hybridization-triggered cleavage agent.

15

20

30

The oligonucleotides may be synthesized by standard methods known in the art, e.g. by use of an automated DNA synthesizer (such as are commercially available from Biosearch, Applied Biosystems, etc.). As examples,

phosphorothioate oligonucleotides may be synthesized by the method of Stein et al. (*Nucl. Acids Res.* 16:3209 (1988)), methylphosphonate oligonucleotides can be prepared by use of controlled pore glass polymer supports (Sarin et al., *Proc. Natl. Acad. Sci. U.S.A.* 85:7448-7451 (1988)), etc.

In a specific embodiment, the MTSP protein antisense oligonucleotide includes catalytic RNA, or a ribozyme (see, *e.g.*, PCT International Publication WO 90/11364, published October 4, 1990; Sarver et al., *Science* 247:1222-1225 (1990)). In another embodiment, the oligonucleotide is a 2'-0-

methylribonucleotide (Inoue et al., *Nucl. Acids Res.* <u>15</u>:6131-6148 (1987)), or a chimeric RNA-DNA analogue (Inoue et al., *FEBS Lett.* <u>215</u>:327-330 (1987)).

5

10

15

20

25

30

In an alternative embodiment, the MTSP protein antisense nucleic acid is produced intracellularly by transcription from an exogenous sequence. For example, a vector can be introduced in vivo such that it is taken up by a cell, within which cell the vector or a portion thereof is transcribed, producing an antisense nucleic acid (RNA). Such a vector would contain a sequence encoding the MTSP protein antisense nucleic acid. Such a vector can remain episomal or become chromosomally integrated, as long as it can be transcribed to produce the desired antisense RNA. Such vectors can be constructed by recombinant DNA technology methods standard in the art. Vectors can be plasmid, viral, or others known in the art, used for replication and expression in mammalian cells. Expression of the sequence encoding the MTSP protein antisense RNA can be by any promoter known in the art to act in mammalian, preferably human, cells. Such promoters can be inducible or constitutive. Such promoters include but are not limited to: the SV40 early promoter region (Bernoist and Chambon, Nature 290:304-310 (1981), the promoter contained in the 3' long terminal repeat of Rous sarcoma virus (Yamamoto et al., Cell 22:787-797 (1980), the herpes thymidine kinase promoter (Wagner et al., Proc. Natl. Acad. Sci. U.S.A. 78:1441-1445 (1981), the regulatory sequences of the metallothionein gene (Brinster et al., Nature 296:39-42 (1982), etc.

The antisense nucleic acids include sequence complementary to at least a portion of an RNA transcript of an MTSP protein gene, preferably a human MTSP protein gene. Absolute complementarily, although preferred, is not required.

The amount of MTSP protein antisense nucleic acid that will be effective in the treatment or prevention of neoplastic disease will depend on the nature of the disease, and can be determined empirically by standard clinical techniques. Where possible, it is desirable to determine the antisense cytotoxicity in cells *in vitro*, and then in useful animal model systems prior to testing and use in humans.

-137-

# 2. Gene Therapy

25

30

In an exemplary embodiment, nucleic acids that include a sequence of nucleotides encoding an MTSP protein or functional domains or derivative thereof, are administered to promote MTSP protein function, by way of gene 5 therapy. Gene therapy refers to therapy performed by the administration of a nucleic acid to a subject. In this embodiment, the nucleic acid produces its encoded protein that mediates a therapeutic effect by promoting MTSP protein function. Any of the methods for gene therapy available in the art can be used (see, Goldspiel et al., Clinical Pharmacy 12:488-505 (1993); Wu and Wu, Biotherapy 3:87-95 (1991); Tolstoshev, An. Rev. Pharmacol. Toxicol. 32:573-10 596 (1993); Mulligan, Science 260:926-932 (1993); and Morgan and Anderson, An. Rev. Biochem. <u>62</u>:191-217 (1993); TIBTECH <u>11(5)</u>:155-215 (1993). For example, one therapeutic composition for gene therapy includes an MTSP protein-encoding nucleic acid that is part of an expression vector that expresses 15 an MTSP protein or domain, fragment or chimeric protein thereof in a suitable host. In particular, such a nucleic acid has a promoter operably linked to the MTSP protein coding region, the promoter being inducible or constitutive, and, optionally, tissue-specific. In another particular embodiment, a nucleic acid molecule is used in which the MTSP protein coding sequences and any other 20 desired sequences are flanked by regions that promote homologous recombination at a desired site in the genome, thus providing for intrachromosomal expression of the MTSP protein nucleic acid (Koller and Smithies, Proc. Natl. Acad. Sci. USA 86:8932-8935 (1989); Zijlstra et al., Nature 342:435-438 (1989)).

Delivery of the nucleic acid into a patient may be either direct, in which case the patient is directly exposed to the nucleic acid or nucleic acid-carrying vector, or indirect, in which case, cells are first transformed with the nucleic acid in vitro, then transplanted into the patient. These two approaches are known, respectively, as in vivo or ex vivo gene therapy.

In a specific embodiment, the nucleic acid is directly administered *in vivo*, where it is expressed to produce the encoded product. This can be accomplished by any of numerous methods known in the art, *e.g.*, by

5

10

15

25

30

-138-

constructing it as part of an appropriate nucleic acid expression vector and administering it so that it becomes intracellular, e.g., by infection using a defective or attenuated retroviral or other viral vector (see U.S. Patent No. 4,980,286), or by direct injection of naked DNA, or by use of microparticle bombardment (e.g., a gene gun; Biolistic, Dupont), or coating with lipids or cellsurface receptors or transfecting agents, encapsulation in liposomes, microparticles, or microcapsules, or by administering it in linkage to a peptide which is known to enter the nucleus, by administering it in linkage to a ligand subject to receptor-mediated endocytosis (see e.g., Wu and Wu, J. Biol. Chem. 262:4429-4432 (1987)) (which can be used to target cell types specifically expressing the receptors), etc. In another embodiment, a nucleic acid-ligand complex can be formed in which the ligand is a fusogenic viral peptide to disrupt endosomes, allowing the nucleic acid to avoid lysosomal degradation. In yet another embodiment, the nucleic acid can be targeted in vivo for cell specific uptake and expression, by targeting a specific receptor (see, e.g., PCT Publications WO 92/06180 dated April 16, 1992 (Wu et al.); WO 92/22635 dated December 23, 1992 (Wilson et al.); WO92/20316 dated November 26, 1992 (Findeis et al.); WO93/14188 dated July 22, 1993 (Clarke et al.), WO 93/20221 dated October 14, 1993 (Young)). Alternatively, the nucleic acid can 20 be introduced intracellularly and incorporated within host cell DNA for expression, by homologous recombination (Koller and Smithies, Proc. Natl. Acad. Sci. USA 86:8932-8935 (1989); Zijlstra et al., Nature 342:435-438 (1989)).

In a specific embodiment, a viral vector that contains the MTSP protein nucleic acid is used. For example, a retroviral vector can be used (see Miller et al., Meth. Enzymol. 217:581-599 (1993)). These retroviral vectors have been modified to delete retroviral sequences that are not necessary for packaging of the viral genome and integration into host cell DNA. The MTSP protein nucleic acid to be used in gene therapy is cloned into the vector, which facilitates delivery of the gene into a patient. More detail about retroviral vectors can be found in Boesen et al., Biotherapy 6:291-302 (1994), which describes the use of a retroviral vector to deliver the mdr1 gene to hematopoietic stem cells in order to make the stem cells more resistant to chemotherapy. Other references

illustrating the use of retroviral vectors in gene therapy are: Clowes et al., *J. Clin. Invest.* 93:644-651 (1994); Kiem et al., *Blood* 83:1467-1473 (1994); Salmons and Gunzberg, *Human Gene Therapy* 4:129-141 (1993); and Grossman and Wilson, *Curr. Opin. in Genetics and Devel.* 3:110-114 (1993).

5

10

15

20

25

30

Adenoviruses are other viral vectors that can be used in gene therapy. Adenoviruses are especially attractive vehicles for delivering genes to respiratory epithelia. Adenoviruses naturally infect respiratory epithelia where they cause a mild disease. Other targets for adenovirus-based delivery systems are liver, the central nervous system, endothelial cells, and muscle. Adenoviruses have the advantage of being capable of infecting non-dividing cells. Kozarsky and Wilson, Current Opinion in Genetics and Development 3:499-503 (1993) present a review of adenovirus-based gene therapy. Bout et al., Human Gene Therapy 5:3-10 (1994) demonstrated the use of adenovirus vectors to transfer genes to the respiratory epithelia of rhesus monkeys. Other instances of the use of adenoviruses in gene therapy can be found in Rosenfeld et al., Science 252:431-434 (1991); Rosenfeld et al., Cell 68:143-155 (1992); and Mastrangeli et al., J. Clin. Invest. 91:225-234 (1993).

Adeno-associated virus (AAV) has also been proposed for use in gene therapy (Walsh et al., *Proc. Soc. Exp. Biol. Med.* 204:289-300 (1993).

Another approach to gene therapy involves transferring a gene to cells in tissue culture by such methods as electroporation, lipofection, calcium phosphate mediated transfection, or viral infection. Usually, the method of transfer includes the transfer of a selectable marker to the cells. The cells are then placed under selection to isolate those cells that have taken up and are expressing the transferred gene. Those cells are then delivered to a patient.

In this embodiment, the nucleic acid is introduced into a cell prior to administration *in vivo* of the resulting recombinant cell. Such introduction can be carried out by any method known in the art, including but not limited to transfection, electroporation, microinjection, infection with a viral or bacteriophage vector containing the nucleic acid sequences, cell fusion, chromosome-mediated gene transfer, microcell-mediated gene transfer, spheroplast fusion, etc. Numerous techniques are known in the art for the

introduction of foreign genes into cells (see *e.g.*, Loeffler and Behr, *Meth. Enzymol.* 217:599-618 (1993); Cohen et al., *Meth. Enzymol.* 217:618-644 (1993); Cline, *Pharmac. Ther.* 29:69-92 (1985)) and may be used, provided that the necessary developmental and physiological functions of the recipient cells are not disrupted. The technique should provide for the stable transfer of the nucleic acid to the cell, so that the nucleic acid is expressible by the cell and preferably heritable and expressible by its cell progeny.

The resulting recombinant cells can be delivered to a patient by various methods known in the art. In a preferred embodiment, epithelial cells are injected, *e.g.*, subcutaneously. In another embodiment, recombinant skin cells may be applied as a skin graft onto the patient. Recombinant blood cells (*e.g.*, hematopoietic stem or progenitor cells) are preferably administered intravenously. The amount of cells envisioned for use depends on the desired effect, patient state, etc., and can be determined by one skilled in the art.

10

15

20

25

30

Cells into which a nucleic acid can be introduced for purposes of gene therapy encompass any desired, available cell type, and include but are not limited to epithelial cells, endothelial cells, keratinocytes, fibroblasts, muscle cells, hepatocytes; blood cells such as T lymphocytes, B lymphocytes, monocytes, macrophages, neutrophils, eosinophils, megakaryocytes, granulocytes; various stem or progenitor cells, in particular hematopoietic stem or progenitor cells, e.g., as obtained from bone marrow, umbilical cord blood, peripheral blood, fetal liver, etc.

In a preferred embodiment, the cell used for gene therapy is autologous to the patient. In an embodiment in which recombinant cells are used in gene therapy, an MTSP protein nucleic acid is introduced into the cells such that it is expressible by the cells or their progeny, and the recombinant cells are then administered *in vivo* for therapeutic effect. In a specific embodiment, stem or progenitor cells are used. Any stem and/or progenitor cells which can be isolated and maintained *in vitro* can potentially be used in accordance with this embodiment. Such stem cells include but are not limited to hematopoietic stem cells (HSC), stem cells of epithelial tissues such as the skin and the lining of the gut, embryonic heart muscle cells, liver stem cells (PCT Publication WO

-141-

94/08598, dated April 28, 1994), and neural stem cells (Stemple and Anderson, *Cell* 71:973-985 (1992)).

Epithelial stem cells (ESCs) or keratinocytes can be obtained from tissues such as the skin and the lining of the gut by known procedures (Rheinwald, *Meth. Cell Bio.* 21A:229 (1980)). In stratified epithelial tissue such as the skin, renewal occurs by mitosis of stem cells within the germinal layer, the layer closest to the basal lamina. Stem cells within the lining of the gut provide for a rapid renewal rate of this tissue. ESCs or keratinocytes obtained from the skin or lining of the gut of a patient or donor can be grown in tissue culture (Rheinwald, *Meth. Cell Bio.* 21A:229 (1980); Pittelkow and Scott, *Mayo Clinic Proc.* 61:771 (1986)). If the ESCs are provided by a donor, a method for suppression of host versus graft reactivity (*e.g.*, irradiation, drug or antibody administration to promote moderate immunosuppression) can also be used.

10

With respect to hematopoietic stem cells (HSC), any technique which 15 provides for the isolation, propagation, and maintenance in vitro of HSC can be used in this embodiment. Techniques by which this may be accomplished include (a) the isolation and establishment of HSC cultures from bone marrow cells isolated from the future host, or a donor, or (b) the use of previously established long-term HSC cultures, which may be allogeneic or xenogeneic. 20 Non-autologous HSC are used preferably in conjunction with a method of suppressing transplantation immune reactions of the future host/patient. In a particular embodiment, human bone marrow cells can be obtained from the posterior iliac crest by needle aspiration (see, e.g., Kodo et al., J. Clin. Invest. 73:1377-1384 (1984)). In a preferred embodiment, the HSCs can be made 25 highly enriched or in substantially pure form. This enrichment can be accomplished before, during, or after long-term culturing, and can be done by any techniques known in the art. Long-term cultures of bone marrow cells can be established and maintained by using, for example, modified Dexter cell culture techniques (Dexter et al., J. Cell Physiol. 91:335 (1977) or Witlock-Witte culture 30 techniques (Witlock and Witte, Proc. Natl. Acad. Sci. USA 79:3608-3612 (1982)).

-142-

In a specific embodiment, the nucleic acid to be introduced for purposes of gene therapy includes an inducible promoter operably linked to the coding region, such that expression of the nucleic acid is controllable by controlling the presence or absence of the appropriate inducer of transcription.

# 3. Prodrugs

5

10

15

20

25

30

A method for treating tumors is provided. The method is practiced by administering a prodrug that is specifically cleaved by an MTSP to release an active drug. Upon contact with a cell that expresses MTSP activity, the prodrug is converted into an active drug. The prodrug can be a conjugate that contains the active agent, such as an anti-tumor drug, such as a cytotoxic agent, or other atherapeutic agent, linked, linked to a substrate for the targeted MTSP, such that the drug or agent is inactive or unable to enter a cell, in the conjugate, but is activated upon cleavage. The prodrug, for example, can contain an oligopeptide, preferably a relatively short, less than about 10 amino acids peptide, that is selectively proteolytically cleaved by the targeted MTSP. Cytotoxic agents, include, but are not limited to, alkylating agents, antiproliferative agents and tubulin binding agents. Others include, vinca drugs, mitomycins, bleomycins and taxanes.

## M. ANIMAL MODELS

Transgenic animal models are provided herein. Such an animal can be initially produced by promoting homologous recombination between an MTSP protein gene in its chromosome and an exogenous MTSP protein gene that has been rendered biologically inactive (preferably by insertion of a heterologous sequence, *e.g.*, an antibiotic resistance gene). In a preferred aspect, this homologous recombination is carried out by transforming embryo-derived stem (ES) cells with a vector containing the insertionally inactivated MTSP protein gene, such that homologous recombination occurs, followed by injecting the ES cells into a blastocyst, and implanting the blastocyst into a foster mother, followed by the birth of the chimeric animal ("knockout animal") in which an MTSP protein gene has been inactivated (see Capecchi, *Science* 244:1288-1292 (1989)). The chimeric animal can be bred to produce additional knockout animals. Such animals can be mice, hamsters, sheep, pigs, cattle, etc., and are

-143-

preferably non-human mammals. In a specific embodiment, a knockout mouse is produced.

Such knockout animals are expected to develop or be predisposed to developing neoplastic diseases and thus can have use as animal models of such diseases e.g., to screen for or test molecules for the ability to treat or prevent such diseases or disorders. Hence, the animal models for are provided. Such an animal can be initially produced by promoting homologous recombination between an MTSP gene in its chromosome and an exogenous MTSP protein gene that would be over-expressed or mis-expressed (preferably by expression under a strong promoter). In a preferred aspect, this homologous recombination is carried out by transforming embryo-derived stem (ES) cells with a vector containing the over-expressed or mis-expressed MTSP protein gene, such that homologous recombination occurs, followed by injecting the ES cells into a blastocyst, and implanting the blastocyst into a foster mother, followed by the birth of the chimeric animal in which an MTSP gene has been over-expressed or mis-expressed (see Capecchi, Science 244:1288-1292 (1989)). The chimeric animal can be bred to produce additional animals with over-expressed or misexpressed MTSP protein. Such animals can be mice, hamsters, sheep, pigs, cattle, etc., and are preferably non-human mammals. In a specific embodiment, a mouse with over-expressed or mis-expressed MTSP protein is produced.

The following examples are included for illustrative purposes only and are not intended to limit the scope of the invention.

**EXAMPLE 1** 

Cloning of MTSP3, cloning and mutagenesis of the Protease domain of MTSP3

- 1. Identification and cloning of MTSP3
  - a. Identification of EST clones Al924527 and Al924182 as part of a serine protease MTSP3

30

25

5

10

15

20

DNA encoding the protease domain of the protease designated MTSP1 was independently cloned from the human prostatic adenocarcinoma cell line, PC-3, using degenerate oligonucleotide primers, then sequenced and

-144-

characterized (see EXAMPLE 6). The sequence of the sense degenerate primer used in cloning MTSP1 was 5'-TGGRT(I)VT(I)WS(I)GC(I)RC(I)CAYTG-3' (SEQ ID No: 13), and that of the anti-sense was

5'-(I)GG(I)CC(I)SWRTC(I)CCYT(I)RCA(I)GHRTC-3' (SEQ ID No:14), where R=A,G; V=G,A,C; W=A,T; S=G,C; Y=C,T; H=A,T,C. The primer sequences correspond to two highly conserved regions in all serine proteases and should amplify PCR products ranging from 400 to 500 base pairs. MTSP1 was subsequently found to be identical to matriptase (Genbank accession number AF118224; see also Takeuchi et al., *Proc. Natl. Acad. Sci. USA*,

10 <u>96(20)</u>:11054-61 (1999); and Lin et al., *J. Biol. Chem.*, <u>274(26)</u>:18231-6 1999).

15

20

25

30

Using the protein sequence of the protease domain of the serine protease MTSP1, the EST database (dbEST) at the National Center for Biotechnology Information (Bethesda, MD; www.ncbi.nlm.nih.gov) was searched for EST clones that contain similar or identical sequences to MTSP1 using the search algorithm tblastn. The tblastn algorithm compares a protein query sequence against a nucleotide sequence database dynamically translated in all six reading frames (both strands). The sequences for two identical EST clones (NCI\_CGAP\_Lu19 Al924527 and Al924182) derived from human lung tumor tissue showed 43% identity with the MTSP1 protein sequence. Subsequent search of GenBank and SwissProt database for the EST sequence Al924527 and Al924182 did not show any matching sequence to MTSP1, indicating that the sequence contained in these EST clones Al924527 and Al924182 may be portions of a new serine protease.

## b. PCR cloning of a cDNA fragment of another membrane type serine protease MTSP3

The double-stranded Marathon-Ready(tm) cDNA library derived from human lung carcinoma (LX-1) was obtained from Clontech (Palo Alto, CA; catalog # 7495-1) and used as a template. Two primers, 5'-TCACCGAGAAGATGATGTGTGCAGGCATCC-3' (SEQ ID No:15) (sense primer), and 5'-GGGACAGGGGCTGTAAGGCAGGGAATGAG-3' (SEQ ID No:16) (antisense primer), were used to amplify a ~360 bp DNA fragment. The PCR

-145-

product was separated on a 2% agarose gel and purified using a gel extraction kit (catalog number 28706; QIAquick gel extraction kit; Qiagen). The purified DNA fragment was ligated into TA vectors (catalog number K4500-01; TOPO-TA cloning kit, Invitrogen, Carlsbad, CA). After transformation into E. coli cells, plasmids were isolated and analyzed by digestion with EcoRI restriction enzyme. Clones that had inserted DNA were further characterized by sequencing using a fluorescent dye-based DNA sequencing method (catalog number 4303149; BigDye terminator cycle sequencing kit with AmpliTaq DNA polymerase; Perkin Elmer, Lincoln, CA).

The DNA sequence obtained was analyzed and has 43% identity with the MTSP1 protein sequence. This indicates that the LX-1 cDNA library contains a desired nucleic acid molecule. It was used to isolate a cDNA clone encompassing a full length protease.

#### c. 5'- and 3'- rapid amplification of cDNA ends (RACE)

To obtain the full-length cDNA that encoded this serine protease, hereafter called MTSP3, 5'- and 3'-RACE reactions were performed. The Marathon-Ready cDNA library from human lung carcinoma (LX-1) was used to isolate the 5' and 3' ends of the cDNA encoding MTSP3. Marathon-Ready cDNA is specifically made for RACE reactions. Two gene specific primers were used: 5'-CCCGCAGCCATAGCCCCAGCTAACG-3' (SEQ ID No. 17) for 5'-RACE reaction and 5'-GCAGACGATGCGTACCAGGGGGAAGTC-3' (SEQ ID No. 18) for 3'-RACE reaction. Two fragments, approximately 1.8 kbp and 0.85 kbp, were isolated that correspond to the missing 5' and 3' end sequences, respectively. These fragments were subcloned as described above. They were further confirmed by Southern analysis using an internal cDNA fragment encompassing the 2 primers used in the RACE reactions as probe and by DNA sequence analysis.

## d. PCR amplification of cDNA encoding full-length protease domain of MTSP3

30

25

10

15

20

To obtain the cDNA fragment encoding the protease domain of MTSP3, an end-to-end PCR amplification using gene-specific primers was used. The two primers used were:

-146-

5'-CTCGAGAAAAGAGTGGTGGGTGGGGAGGAGGCCTCTGTG-3' (SEQ ID No. 19) for the 5' end and 5'-GCGGCCGCATTACAGCTCAGCCTTCCAGAC-3' (SEQ ID No. 20) for the 3' end. The 5' primer contains the sequence (underlined) that encodes the start of the MTSP3 protease domain (VVGGEEASV). The 3' primer contains the stop codon (underlined) of MTSP3. A ~700-bp fragment was amplified and subcloned into a Pichia pastoris expression vector, pPIC9K.

### e. C310S mutagenesis of MTSP3

To eliminate the free cysteine (at position 310 in SEQ ID No. 4) that exists when the protease domain of the MTSP3 protein is expressed or the zymogen is activated, the free cysteine at position 310 (see SEQ ID No. 3), which is Cys122 if a chymotrypsin numbering scheme is used, was replaced with a serine. The resulting vector was designated pPIC9K:MTSP3C122S.

The gene encoding the protease domain of MTSP3 was mutagenized by PCR SOE (PCR-based splicing by overlap extension) to replace the unpaired cysteine at position 310 (122 chymotrypsin numbering system) with a serine. Two overlapping gene fragments, each containing the TCT codon for serine at position 310 were PCR amplified using the following primers: for the 5' gene fragment, TCTCTCGAGAAAAGAGTGGTGGGTGGGTGGGGAGGAGGCCTCTGTG SEQ ID No. 51 and

20 GCTCCTCATCAAAGAAGGGCAGAGAGATGGGCCTGACTGTGCC SEQ ID No. 52; for the 3' gene fragment,

ATTCGCGGCCGCATTACAGCTCAGCCTTCCAGAC (SEQ ID No. 53) and GGCACAGTCAGGCCCATCTCTCTGCCCTTCTTTGATGAGGAGC (SEQ ID No.

54). The amplified gene fragments were purified on a 1% agarose gel, mixed and reamplified by PCR to produce the full length coding sequence for MTSP3 C122S. This sequence was then cut with restriction enzymes Notl and Xhol, and ligated into vector pPic9K.

#### 2. Sequence analysis

10

15

25

30

All derived DNA and protein sequences were analyzed using MacVector (version 6.5; Oxford Molecular Ltd., Madison, WI). The full-length cDNA encoding MTSP3 is composed of 2,137 base pairs containing the longest open reading frame of 1,314 base pairs which translate to a 437-amino acid protein

-147-

sequence. The cDNA fragment (nt 873-1,574) encoding the protease domain of MTSP3 is composed of 702 base pairs which translate to a 233-amino acid protein sequence plus the stop codon. The DNA sequence and the translated protein sequence of MTSP3 are shown in SEQ ID Nos. 3 and 4, respectively.

#### 5 3. Construction of the Expression Vectors

extension at 75 °C for 2 min.

10

15

20

25

30

DNA encoding MTSP3 full length protein containing the C310S point mutation (i.e., MTSP3C122S) was cloned from pPIC9K:MTSP3C122S. The primers MTSP3:

5' GAATTCCATATGCCGCGCTTTAAAGTGGTGGGTGGGAGGAGGCC SEQ ID No. 47 (containing a Ndel restriction site) and MTSP3-

3' GGGATACCCGTTACAGCTCAGCCTTCCAGAC 5' SEQ ID No. 48 (containing a BamHI restriction site) were used to PCR amplify the human MTSP3C122S protease domain utilizing a plasmin recognition sequence (PRFK) for zymogen activation. Amplification was conducted in a total volume of 50  $\mu$ l containing 10 mM KCI, 20 mM Tris-HCI (pH 8.8 at 25 °C), 10 mM (NH<sub>4</sub> )<sub>2</sub> SO<sub>4</sub> , 2.0 mM MgSO<sub>4</sub> , 0.1% Triton X-100, 0.3 mM dNTPs, 5.0 units of vent DNA polymerase, and 100 pmol of primers. The reaction mixtures were heated to 95 °C for 5 min, followed by 25–30 cycles of 95, 60, and 75 °C for 30 s each and a final

PCR products were purified using a QIAquick PCR purification kit (QIAGEN Inc., Chatsworth, CA). Full-length oligonucleotides were doubly digested with 10 units BamHI and 20 units Ndel for 2 h at 37°C. The digested fragments were purified on a 1.3% agarose gel and stained with ethidium bromide. The band containing the MTSP3C122S encoding DNA was excised and purified using a QIAEX II gel extraction kit.

The MTSP3C122S encoding DNA was then cloned into the Ndel and BamHI sites of the pET19b vector (Novagen) using standard methods. This vector allows the fusion of a HIS<sub>6</sub> tag for purification by metal affinity chromatography (MAC). Competent XL1 Blue cells (Stratagene) were transformed with the pET19b-MTSP3C122S vector and used to produce plasmid stocks. Proper insertion and DNA sequence were confirmed by fluorescent thermal dye DNA sequencing methods as well as restriction digests.

-148-

### 4. Protein Expression, Purification, and Refolding

10

15

20

25

30

Overexpression of the gene product was achieved in *E. coli* strain BL21(DE3) (Novagen, Madison WI) containing the DNAY plasmid for rare codon optimization (see, *e.g.*, Garcia *et al.* (1986) *Cell 45:*.453-459). Cells were grown at 37 °C in (2xYT) media supplemented with carbenicillin and kanamycin to a final concentrations of 50 ug/ml and 34 ug/ml, respectively. One liter cultures were inoculated with 10 mL of an overnight culture grown in the same media. Cells were allowed to grow to a density of 0.6 – 1.0 OD<sub>600</sub> before the addition of IPTG (final concentration 1.0 mM). Cells were grown an additional 4 hours before harvesting.

The cell pellet was resuspended in 20 mL of lysis buffer (50 mM  $\rm Na_2HPO_4$ , 300 mM NaCl, pH 7.4). The cell suspension was treated with 10-20 mg lysozyme and incubated at 37°C for 1 hour. DNasel was then added (1-2mg) with mixing until the solution was no longer viscous. The solution was then transferred to a Rosette flask and sonicated, on ice, at high power for 15 min. Inclusion bodies were pelleted by centrifugation at 20K rpm ( $\sim$ 48,000 g) at 4°C for 30 min.

Inclusion bodies were washed by douncing 2 times in 50 mM  $\rm Na_2HPO_4$ , 300 mM  $\rm NaCl$ , 5% LADO, pH 7.4 followed by 2 times in 50 mM  $\rm Na_2HPO_4$ , 300 mM  $\rm NaCl$ , pH 7.4. Inclusion bodies ( $\sim$  500 mg) are solubilized in 25 mL 6 M GuHCl, 100 mM tris-HCl, 20 mM  $\beta$ Me, pH 8.0. This solution was spun at 20K rpm for 30 minutes to pull down any particulate matter. This solution was passed through a 0.2  $\mu$ M filter and diluted to 100 mL in solubilization buffer.

MTSP3C122S was refolded by slowly adding the inclusion body mixture to 8 L of refolding buffer (100 mM tris-HCl, 150 mM NaCl, 5 mM GSH, 0.05 mM GSSG, 1 M arginine, pH 8.0) using a peristaltic pump. The refolding mixture was allowed to stir at  $4^{\circ}$ C for 7 days or until the thiol concentration was below 1 mM as detected by Ellman's reagent. The solution was filtered through a 5  $\mu$ M filter, concentrated by ultrafiltration and the buffer exchanged into MAC equilibration buffer (50 mM Na<sub>2</sub>HPO<sub>4</sub>, 300 mM NaCl, 10 mM imidazole, pH 8.0) by crossflow filtration. The resulting solution was passed through a 0.2  $\mu$ M filter and further purified on a FPLC (Amersham-Pharmacia)

-149-

using Pharmacia chelating sepharose. The solution was loaded onto the nickel loaded MAC at a flow rate of 1.0 mL/min and eluted with a linear gradient of 1.0 mM -1.0 M imidazole in 50 mM  $Na_2HPO_4$ , 300 mM NaCl, pH 8.0. Protein containing fractions were determined by SDS-PAGE and subsequently pooled and frozen at  $-80\,^{\circ}C$ .

Small amounts of purified MTSP3C122S were activated using plasmin sepharose for 30 min. at 37°C. The resin was spun down at 14K rpm for 5 min. and the protein solution removed. The resulting solution was screened for activity against of series of protease substrates; spec-tpa, spec-pl, spec-UK, spec-fXIIa (American Diagnostica), S-2238, S-2266 (Kabi Diagnostica), S-2586, S-2366, S-2444, S-2288, S-2251, S-2302, S-2765, S-2222, spec-THE (Chromogenix), spec-fVIIa (Pentapharm). MTSP3C122S cleaved several of these substrates efficiently but was most active towards Spec-fXIIa, Spec-tPA, S-2765, Spec-fVIIa and S-2444.

### 15 5. Gene expression profile of the serine protease MTSP3 in normal and tumor tissues

20

25

30

To obtain information regarding the tissue distribution of the MTSP3 transcripts, the DNA insert encoding the MTSP3 protease domain was used to probe a RNA blot composed of 76 different human tissues (catalog number 7775-1; human multiple tissue expression (MTE) array; CLONTECH, Palo Alto, CA). The expression pattern observed in decreasing signal level was: trachea = colon (descending) = esophagus > colon (ascending) > colon (transverse) = rectum > ileum > duodenum > jejunum > bladder > ilocecum > stomach > kidney > appendix. It is also expressed less abundantly in fetal kidney, and in two tumor cell lines, HeLa S3 and leukemia, K-562. Northern analysis using RNA blots (catalog numbers 7780-1, 7765-1 & 7782-1; human 12-lane, human muscle and human digestive system multiple tissue northern (MTN) blots; CLONTECH) confirmed that the expression was detected most abundantly in the colon, moderately in the esophagus, small intestine, bladder and kidney, and less abundantly in stomach and rectum. A single transcript of ~2.2 kb was detected.

-150-

Amplification of the MTSP3 transcript in several human primary tumors xenografted in mouse was performed using gene-specific primers. The MTSP3 transcript was detected in lung carcinoma (LX-1), colon adenocarcinoma (CX-1), colon adenocarcinoma (GI-112) and ovarian carcinoma (GI-102). No apparent signal was detected in another form of lung carcinoma (GI-117), breast carcinoma (GI-101), pancreatic adenocarcinoma (GI-103) and prostatic adenocarcinoma (PC3).

#### **EXAMPLE 2**

15

#### Identification of genomic clone of MTSP4

10

15

20

25

30

Using the nucleotide sequence encoding the protease domain of the serine protease MTSP1 (also called matriptase), the protein database (SWISSPROT) at the National Center for Biotechnology Information (Bethesda, MD; <a href="http://www.ncbi.nlm.nih.gov">http://www.ncbi.nlm.nih.gov</a>) was searched for similar or identical sequence to MTSP1 using the search algorithm blastx. The blastx algorithm compares the six-frame conceptual translation products of a nucleotide query sequence (both strands) against a protein sequence database. A protein encoding sequence (CAA18442) that has 37% identity to the MTSP1 protein sequence that was found to include a putative LDL-receptor domain and a trypsin-like serine protease domain was identified. This protein-encoding sequence (hereinafter referred to as MTSP4) was found to be encoded by a genomic clone (AL022314) derived from human chromosome 22 sequenced by the Sanger Centre Chromosome 22 Mapping Group and deposited into the public database as part of the Human Genome Project. Subsequent search of the GenBank database showed that no identical sequence has been deposited. A search of the EST database also did not show any matching human sequence, indicating that no human EST clone exists in the public database. Mouse EST clones (Al391417 and AA208793) are present and showed 88% identity to the serine protease at the nucleotide level.

## PCR cloning of a genomic DNA fragment of MTSP4 for use as hybridization probe

In order to obtain tissue distribution profile of MTSP4 as well as to identify a tissue source for subsequent cloning of the cDNA, a genomic fragment

-151-

was amplified from human genomic DNA using two gene-specific primers, 5'-CCTCCACGGTGCTGTGGACCGTGTTCC-3' (5' primer) SEQ ID No. 21 and 5'-CCTCGCGCAAGGCGCCCCAGCCCG-3' (3' primer) SEQ ID No. 22. These two primers amplified a 265-base pair fragment within a single exon of MTSP4.

5 The fragment was then used as a hybridization probe on human tissue northern blot (human 12-lane multiple tissue northern (MTN) blot (catalog number 7780-1); CLONTECH, Palo Alto, CA). A prominent band (~2.6 kb) was detected in liver. Relatively weaker signals were obtained from the brain, heart, skeletal muscle and kidney. Since human liver showed a very strong signal, this tissue was selected for the amplification of the MTSP4 cDNA.

#### 5'- and 3'- rapid amplification of cDNA ends (RACE)

15

20

25

30

To obtain a full-length clone encoding MTSP4, 5'- and 3'-RACE reactions were performed. The Marathon-Ready cDNA library from human liver (CLONTECH) was used to isolate the 5' and 3' ends of the cDNA encoding MTSP4. Marathon-Ready cDNA clones are specifically made for RACE reactions. Two gene specific primers were used: 5'-GCGTGGCGTCACCTGGTAGCGATAGACCTCGC -3' (SEQ ID No. 23) for 5'-RACE reaction and 5'-CCTCCACGGTGCTGTGGACCGTGTTCC-3' (SEQ ID No. 24) for 3'-RACE reaction. No fragment was obtained from the initial 5'-RACE reaction.

The 3'-RACE reaction, however, produced a ~1.5 kbp fragment. A nested PCR reaction was used on the initial 5'-RACE reaction products to obtain part of the 5' end of MTSP4. The nested 5' gene-specific primer used was 5'-CCTCGCGCAAGGCGCCCCAGCCCG-3' (SEQ ID No. 25) and produced a ~0.8 kbp fragment. The fragments were subcloned into pCR2.1-TOPO TA cloning vector (Invitrogen, Carlsbad, CA). The resulting clones were analyzed by Southern analysis using the internal genomic fragment encompassing the primers used in the RACE reactions as probe and by DNA sequence analysis. Sequence analysis of the 5'-RACE product showed that the potential initiation codon was still missing.

To obtain the 5' cDNA end that encodes the N terminus of MTSP4, the publicly available genomic sequence of chromosome 22 was searched for

-152-

sequence corresponding to the sequence obtained in the 5'-RACE clone. The resulting genomic sequence was translated and the protein sequence was compared to that derived from the translated sequence of the 5'-RACE clone. After determining the overlapping sequences, a gene-specific oligonucleotide primer (5'-TCATCGGCCAGAGGGTGATCAGTGAG-3') SEQ ID No. 26 corresponding to the sequence upstream of the potential initiation codon and another gene-specific oligonucleotide primer (5'-CCTCCTCAGTGCATAGGCATCAAACCAG-3') SEQ ID No. 27 corresponding to a sequence within the overlapping region were used to amplify the missing 5' cDNA of MTSP4 from the human liver cDNA library.

### Splice variants and domain organization of MTSP4

5

10

15

20

25

30

At least two cDNA fragments were consistently obtained during PCR amplification, indicating multiple splice variants of MTSP4. Subcloning and sequence analysis revealed that a longer, more abundant form, MTSP4-L and a shorter form, MTSP4-S. The encoded proteins are multi-domain, type II membrane-type serine proteases and include a transmembrane domain at the N terminus followed by a CUB domain, 3 LDLR domains and a trypsin-like serine protease domain at the C terminus. The difference between these two forms of MTSP4 is the absence in MTSP4-S of a 432-bp nucleotide sequence between the transmembrane and the CUB domains (see FIGURE 2).

### PCR amplification of cDNA encoding full-length protease domain of MTSP4

To obtain a cDNA fragment encoding the protease domain of MTSP4, an end-to-end PCR amplification using gene-specific primers and the Marathon-Ready cDNA library from human liver was used. The two primers used were: 5'-TCTCTCGAGAAAAGAATTGTTGGTGGAGCTGTGTCCTCCGAG-3' (SEQ ID No. 28) for the 5' end and 5'-AGGTGGGCCTTGCTTTGCAGGGGGGCAGTTC-3' for the 3' end SEQ ID NO. 29). The 5' primer contained the sequence that encodes the start of the MTSP4 protease domain (IVGGAVSSE). The 3' primer corresponds to the sequence just downstream of the stop codon. A ~740-bp fragment was amplified, subcloned into pCR2.1-TOPO TA cloning vector and sequenced.

Gene expression profile of MTSP4 in normal and tumor tissues

-153-

To obtain information regarding the gene expression profile of the MTSP4 transcript, a DNA fragment encoding part of the LDL receptor domain and the protease domain was used to probe an RNA blot composed of 76 different human tissues (catalog number 7775-1; human multiple tissue expression (MTE) array; CLONTECH). As in the northern analysis of gel blot, a very strong signal was observed in the liver. Signals in other tissues were observed in (decreasing signal level): fetal liver > heart = kidney = adrenal gland = testis = fetal heart and kidney = skeletal muscle = bladder = placenta > brain = spinal cord = colon = stomach = spleen = lymph node = bone marrow = trachea = uterus = pancreas = salivary gland = mammary gland = lung. MTSP4 is also expressed less abundantly in several tumor cell lines including HeLa S3 = leukemia K-562 = Burkitt's lymphomas (Raji and Daudi) = colorectal adenocarcinoma (SW480) > lung carcinoma (A549) = leukemia MOLT-4 = leukemia HL-60. PCR of the MTSP4 transcript from cDNA libraries made from several human primary tumors xenografted in nude mice (human tumor multiple tissue cDNA panel, catalog number K1522-1, CLONTECH) was performed using MTSP4-specific primers. The MTSP4 transcript was detected in breast carcinoma (GI-101), lung carcinoma (LX-1), colon adenocarcinoma (GI-112) and pancreatic adenocarcinoma (GI-103). No apparent signal was detected in another form of lung carcinoma (GI-117), colon adenocarcinoma (CX-1), ovarian carcinoma (GI-102). and prostatic adenocarcinoma (PC3). The MTSP4 transcript was also detected in LNCaP and PC-3 prostate cancer cell lines as well as in HT-1080 human fibrosarcoma cell line.

#### Sequence analysis

10

15

20

25

30

MTSP4 DNA and protein sequences were analyzed using MacVector (version 6.5; Oxford Molecular Ltd., Madison, WI). The ORF of MTSP4-L includes 2,409 bp, which translate to a 802-amino acid protein, while the ORF of MTSP4-S is composed of 1,977 bp which translate to a 658-amino acid protein. The cDNA encoding the protease domain in both forms is composed of 708 bp which translate to a 235-amino acid protein sequence (see, SEQ ID No. 6) The DNA sequences and the translated protein sequences of MTSP4-L and

-154-

MTSP4-S, and of the protease domain of MTSP4 are set forth in SEQ ID Nos. 8, 10 and 6, respectively.

#### **EXAMPLE 3**

#### Cloning of MTSP6

5

20

25

30

#### Identification of genomic clone of MTSP6

Using the protein sequence of the protease domain of the serine protease MTSP4 (see EXAMPLE 2), the non-redundant database (all non-redundant GenBank CDS translations + PDB + SwissProt + PIR + PRF) at the National Center for Biotechnology Information (Bethesda, MD;

<http://www.ncbi.nlm.nih.gov>) was searched for sequences that were similar or identical to MTSP4 using the search algorithm tblastn. The tblastn algorithm compares a protein query sequence against a nucleotide sequence database dynamically translated in all reading frames. A protein (55 amino acids), which has 60% identity with the query MTSP4 sequence (55 amino acids), was obtained from the translation of genomic sequence of ACO15555 (nucleotide #15553 to 15717). This protein hereafter is referred to as MTSP6. Subsequent search of the GenBank database showed that no cDNA encoding MTSP6 has been deposited.

The gene exhibiting highest homology to MTSP6 was human transmembrane serine protease 2 (GenBank accession number U75329; Swissprot accession number O15393), which showed 66% identity to MTSP6 within the 45 amino acid regions compared. Consequently, the nucleotide sequence encoding the MTSP6 protease domain was obtained by comparing the protein sequence of human transmembrane serine protease 2 protease domain with the nucleotide sequence of ACO15555 translated in six reading frames. The protein sequence obtained from the translated nucleotide sequence of MTSP6 revealed an overall 50% identity with human transmembrane serine protease 2. A search of the EST database indicated the presence of seven MTSP6 EST clones (AA883068, AW591433, Al978874, Al469095, Al935487, AA534591 and Al758271).

-155-

### Cloning of human MTSP6 full-length cDNA

To obtain cDNA encoding the region of the MTSP6 protease domain identified by database searches described above, two gene-specific primers, Ch17-NSP-1, 5'-TCACGCATCGTGGGTGGAACATGTCC-3' (5' primer) SEQ ID NO. 30 nd Ch17-NSP-2AS, 5'- ACCCACCTCCATCTGCTCGTGGATCC-3' SEQ ID NO. 31 (3' primer), were used for PCR. These two primers amplified a 708-base pair fragment from human mammary gland carcinoma cDNA (Clontech Marathon-Ready cDNA, Cat. No. 7493-1).

To obtain the remaining, unknown cDNA of MTSP6, 5'- and 3'-RACE reactions were performed on the human mammary gland carcinoma. 10 Marathon-Ready cDNA is specifically made for RACE reactions. The first RACE reactions were performed by PCR using Marathon cDNA adaptor primer 1 (AP1) with gene specific primers, Ch17-NSP-2AS, 5'-ACCCACCTCCATCTGCTCGTGGATCC-3' SEQ ID NO. 31 for 5'-RACE reaction and Ch17-NSP-1, 5'-TCACGCATCGTGGGTGGAACATGTCC-3' SEQ ID NO. 30 15 for 3'-RACE reaction. The PCR products were purified from agarose gel. A second nested PCR was then performed using Marathon cDNA adaptor primer 2 (AP2) with gene specific primer Ch17-NSP-3AS, 5'- CCACAGCCTCCTCTTGACACACCAG-3' SEQ ID No. 32 for 5'-RACE reaction (using first 5'-RACE product as template) and Ch17-NSP-3 20 5'-ACGCCCCTGTGGATCATCACTGCTGC-3' SEQ ID No. 33 for 3'-RACE reaction (using first 3'-RACE product as template). First 5'- and 3'-RACE products were also used as template for PCR reactions using primers Ch17-NSP-3 and Ch17-NSP-4AS to obtain a cDNA fragment for use as a probe. PCR products from RACE reactions which were larger than 700 bp were cut out 25 and purified from agarose gel and subcloned into pCR2.1-TOPO cloning vector (Invitrogen, Carlsbad, CA). Colony hybridization was then performed to identify positive colonies containing MTSP6 sequence. Positive clones were identified by colony hybridization using the 495 bp DNA fragment obtained from PCR reaction (with primers Ch17-NSP-3 and Ch17-NSP-4AS) and by DNA 30 sequencing.

-156-

Sequence analysis of the 5'-RACE products indicated that an additional 420 bp of upstream sequence were obtained. The potential initial codon was not present in the 5'-RACE sequence. Another round of nested 5'-RACE reaction was performed using AP2 and a gene specific primer (designed based on the new RACE sequence) Ch17-NSP-5AS 5'-TCCCTCCCTCACATATACTGAGTGGTG-3' SEQ ID No. 34, using the PCR products obtained from the first 5'-RACE as template. A PCR product of 367 bp using Ch17-NSP-6 5'-CGACTGCTCAGGGAAGTCAGATGTCG-3' SEQ ID No. 35 (designed based on the new 5'-RACE sequence) and Ch17-NSP-5AS was used to identify the positive clones. An additional sequence of 480 bp was obtained from the second 5'-RACE products. A potential ATG start codon was observed within a sequence of GTCACCATGG (nucleotides 262-272 of SEQ ID No. 12, which appears to be a Kozak sequence (GCC (A/G) CCAUGG), indicating that this ATG is likely the initiation codon for MTSP6.

The 3'-RACE reaction to obtain the rest of the 3' end of MTSP6 was not successful using Marathon Ready human mammary gland carcinoma cDNA. The sequence of the 3'-RACE products obtained was exclusively that of an MTSP6 cDNA truncated with the Marathon AP2 primer sequence within the coding region.

The 3'-end sequence of MTSP6 was obtained by PCR using Ch17-NSP-3 (5'-ACGCCCTGTGGATCATCACTGCTGC-3'; SEQ ID NO. 33) and Ch17-NSP-4 (5'-CTGGTGTCAAGAGAGGAGGCTGTGG-3'; SEQ ID NO. 37) with an antisense primer Ch17-NSP-7AS (5'-ACTCAGGTGGCTACTTATCCCCTTCCTC-3'; SEQ ID NO. 38) designed based on the sequence of an EST clone AA883068, which apparently covers the

on the sequence of an EST clone AA883068, which apparently covers the 3'-end of MTSP6 sequence, and human small intestine cDNA (Clontech) as template. Two PCR products (650 bp and 182 bp, respectively) were obtained and DNA sequence analysis indicated that both PCR products contained a stop codon.

#### 30 Sequence analysis and domain organization of MTSP6

10

15

The MTSP6 DNA and protein sequences were analyzed using DNA Strider (version 1.2). The ORF of MTSP6 is composed of 1,362 bp, which translate

-157-

into a 453-amino acid protein. Protein sequence analysis using the SMART (Simple Modular Architecture Research Tool) program at http://smart.embl-heidelberg.de predicts that MTSP6 is a multi-domain, type-II membrane-type serine protease containing of a transmembrane domain (amino acids 48-68) at the N terminus followed by a LDLRa domain (LDL receptor domain class a) (amino acids 72-108), a SR domain (Scavenger receptor Cys-rich domain)(amino acids 109-205), and a trypsin-like serine protease domain (amino acids 216-443) (see FIGURE 3).

#### Gene expression profile of MTSP6 in normal and tumor tissues

10

15

20

25

30

To obtain information regarding the gene expression profile of the MTSP6 transcript, a 495 bp DNA fragment obtained from PCR reaction with primers Ch17-NSP-3 and NSP-4AS was used to probe an RNA blot composed of 76 different human tissues (catalog number 7775-1; human multiple tissue expression (MTE) array; CLONTECH). The strongest signal was observed in duodenum. Signal in other tissues were observed in (decreased signal level): Stomach > trachea = mammary gland = thyroid gland = salivary gland = pituitary gland = pancreas > kidney > lung > jejunum = ileum = ilocecum = appendix = fetal kidney > fetal lung. Very weak signals can also be detected in several other tissues. MTSP6 is also expressed in several tumor cell lines including HeLa S3 > colorectal adenocarcinoma (SW480) > leukemia MOLT-4 > leukemia K-562. PCR analysis of the MTSP6 transcript from cDNA libraries made from several human primary tumors xenografted in nude mice (human tumor multiple tissue cDNA panel, catalog number K1522-1, CLONTECH) was performed using MTSP6-specific primers (Ch17-NSP-3 and Ch17-NSP2AS). The MTSP6 transcript was strongly detected in lung carcinoma (LX-1), moderately detected in pancreatic adenocarcinoma (GI-103), weakly detected in ovarian carcinoma (GI-102); and very weakly detected in colon adenocarcinoma (GI-112 and CX-1), breast carcinoma (GI-101), lung carcinoma (GI-117) and prostatic adenocarcinoma (PC3). The MTSP6 transcript was also detected in breast cancer cell line MDA-MB-231, prostate cancer cell line PC-3, but not in HT-1080 human fibrosarcoma cell line. MTSP6 is also expressed in mammary gland carcinoma cDNA (Clontech).

-158-

#### **EXAMPLE 4**

#### Expression of the protease MTSP domains

The DNA encoding each of the MTSP 3 and 4 protease domains was cloned into a derivative of the *Pichia pastoris* vector pPIC9K (available from Invitrogen; see SEQ ID NO. 45). Plasmid pPIC9k features include the 5' AOX1 promoter fragment at 1-948; 5' AOX1 primer site at 855-875; alpha-factor secretion signal(s) at 949-1218; alpha-factor primer site at 1152-1172; multiple cloning site at 1192-1241; 3' AOX1 primer site at 1327-1347; 3' AOX1 transcription termination region at 1253-1586; HIS4 ORF at 4514-1980; kanamycin resistance gene at 5743-4928; 3' AOX1 fragment at 6122-6879; CoIE1 origin at 7961-7288; and the ampicillin resistance gene at 8966-8106.

CoIE1 origin at 7961-7288; and the ampicillin resistance gene at 8966-8106. The plasmid used herein is derived from pPIC9K by eliminating the Xhol site in the kanamycin resistance gene and the resulting vector is herein designated pPIC9KX.

Primers used for PCR amplification of protease domain and subcloning into the Xhol/Notl sites of Pichia vector

MTSP3

- 5' primer (with Xhol site [underlined]) SEQ ID No. 39
- 5' TCTCTCGAGAAAAGAGTGGTGGGTGGGGAGGAGGCCTCTGTG 3'
- 20 3' primer (with Notl site [underlined]) SEQ ID No. 40
  - 5' ATTCGCGGCCGCATTACAGCTCAGCCTTCCAGAC 3'

#### MTSP4-S and MTSP4-L

- 5' primer (with Xhol site [underlined]) SEQ ID No. 41
  - 5' TCTCTCGAGAAAAGAATTGTTGGTGGAGCTGTGTCCTCCGAG
- 25 3' primer with Notl site SEQ ID No. 42
  - 5' ATTC<u>GCGGCCGC</u>TCAGGTCACCACTTGCTGGATCCAG 3'

#### MTSP6

30

10

15

MTSP6 was cloned into the *E. coli* TOPO vector (pcR® 2.1 TOPO™, SEQ ID No. 46, Invitrogen, Carlsbad, CA; the TOPO® TA Cloning® Kit is designed form cloning Taq-amplified PRCR products).

- 5' primer (with Xhol site [underlined]) SEQ ID No. 43
  - 5' CTCGAGAAACGCATCGTGGGTGGAAACATGTCCTTG 3'

-159-

3' primer NotI site comes from E. coli TOPO vector SEQ ID No. 44:

#### 5' ACTCAGGTGGCTACTTATCCCCTTCCTC 3'

#### **EXAMPLE 5**

Assays for identification of candidate compounds that modulate that activity of an MTSP

#### Assay for identifying inhibitors

5

10

15

20

25

30

The ability of test compounds to act as inhibitors of catalytic activity of an MTSP, including MTSP1, MTSP3, MTSP4, MTSP6 can be assessed in an amidolytic assay. The inhibitor-induced inhibition of amidolytic activity by a recombinant MTSP or the protease domain portions thereof, can be measured by IC50 values in such an assay.

An exemplary assay buffer is HBSA (10 mM Hepes, 150mM sodium chloride, pH 7.4, 0.1% bovine serum albumin). All reagents were from Sigma Chemical Co. (St. Louis, MO), unless otherwise indicated. Two IC50 assays at 30-minute (a 30-minute preincubation of test compound and enzyme) and at 0-minutes (no preincubation of test compound and enzyme) are conducted. For the IC50 assay at 30-minute, the following reagents are combined in appropriate wells of a Corning microtiter plate: 50 microliters of HBSA, 50 microliters of the test compound, diluted (covering a broad concentration range) in HBSA (or HBSA alone for uninhibited velocity measurement), and 50 microliters of the MTSP or protease domain thereof diluted in buffer, yielding a final enzyme concentration of about 100-500 pM. Following a 30-minute incubation at ambient temperature, the assay is initiated by the addition of 50 microliters of a substrate for the particular MTSP (see, e.g., table and discussion below) and reconstituted in deionized water, followed by dilution in HBSA prior to the assay) were added to the wells, yielding a final volume of 200 microliters and a final substrate concentration of 300  $\mu M$  (about 1.5-times Km).

For an IC50 assay at 0-minute, the same reagents are combined: 50 microliters of HBSA, 50 microliters of the test compound, diluted (covering the identical concentration range) in HBSA (or HBSA alone for uninhibited velocity measurement), and 50 microliters of the substrate, such as a chromogenic substrate. The assay is initiated by the addition of 50 microliters of MTSP. The

-160-

final concentrations of all components are identical in both IC50 assays (at 30and 0-minute incubations).

The initial velocity of the substrate hydrolysis is measured in both assays by, for example for a chromogenic substrate, as the change of absorbance at a particular wavelength, using a Thermo Max¿ Kinetic Microplate Reader (Molecular Devices) over a 5 minute period, in which less than 5% of the added substrate was used. The concentration of added inhibitor, which caused a 50% decrease in the initial rate of hydrolysis was defined as the respective IC50 value in each of the two assays (30-and 0-minute).

#### Another assay for identifying inhibitors

Test compounds for inhibition of the protease activity of the protease domain of is assayed in Costar 96 well tissue culture plates (Corning NY). Approximately 2-3 nM the MTSP or protease domain thereof is mixed with varying concentrations of inhibitor in 29.2 mM Tris, pH 8.4, 29.2 mM imidazole, 217 mM NaCl (100 mL final volume), and allowed to incubate at room temperature for 30 minutes. 400 mM substrate is added, and the reaction monitored in a SpectraMAX Plus microplate reader (Molecular Devices, Sunnyvale CA) by following the change in a parameter correlated with hydrolysis, such as absorbance for a chromogenic substrate for 1 hour at 37° C.

20

25

30

5

10

15

#### **ASSAY FOR SCREENING MTSP6**

The protease domain of MTSP6 expressed in Pichia pastoris is assayed for inhibition by various compounds in Costar 96 well tissue culture plates (Corning NY). Approximately 1-20 nM MTSP6 is mixed with varying concentrations of inhibitor in 29.2 mM Tris, pH 8.4, 29.2 mM Imidazole, 217 mM NaCl (100  $\mu$ L final volume), and allowed to incubate at room temperature for 30 minutes. 500  $\mu$ M substrate Spectrozyme t-PA (American Diagnostica,, Greenwich, CT) is added, and the reaction is monitored in a SpectraMAX Plus microplate reader (Molecular Devices, Sunnyvale CA) by measuring the change in absorbance at 405 nm for 30 minutes at 37 °C.

#### Identification of substrates

-161-

Particular substrates for use in the assays can be identified empirically by testing substrates. The following list of substrates are exemplary of those that can be tested.

	Substrate name	Structure
5	S 2366	pyroGlu-Pro-Arg-pNA.HCl
	spectrozyme t-PA	CH₃SO₂-D-HHT-Gly-Arg-pNA.AcOH
	N-p-tosyl-Gly-Pro-Arg-pNA	N-p-tosyl-Gly-Pro-Arg-pNA
	Benzoyl-Val-Gly-Arg-pNA	Benzoyl-Val-Gly-Arg-pNA
	Pefachrome t-PA	CH <sub>3</sub> SO <sub>2</sub> -D-HHT-Gly-Arg-pNA
10	S 2765	N-α-Z-D-Arg-Gly-Arg-pNA.2HCl
	S 2444	pyroGlu-Gly-Arg-pNA.HCl
	S 2288	H-D-IIe-Pro-Arg-pNA.2HCI
	spectrozyme UK	Cbo-L-(y)Glu(a-t-BuO)-Gly-Arg-pNA.2AcOH
	S 2302	H-D-Pro-Phe-Arg-pNA.2HCI
15	S 2266	H-D-Val-Leu-Arg-pNA.2HCl
	S 2222	Bz-Ile-Glu(g-OR)-Gly-Arg-pNA.HCl
		$R = H(50\%)$ and $R = CH_3(50\%)$
	Chromozyme PK	Benzoyl-Pro-Phe-Arg-pNA
	S 2238	H-D-Phe-Pip-Arg-pNA.2HCl
	S 2251	H-D-Val-Leu-Lys-pNA.2HCl
20	Spectrozyme PI	H-D-Nle-HHT-Lys-pNA.2AcOH
		Pyr-Arg-Thr-Lys-Arg-AMC
		H-Arg-Gln-Arg-Arg-AMC
		Boc-Gln-Gly-Arg-AMC
		Z-Arg-Arg-AMC
25	Spectrozyme THE	H-D-HHT-Ala-Arg-pNA.2AcOH
	Spectrozyme fXIIa	H-D-CHT-Gly-Arg-pNA.2AcOH
		CVS 2081-6 (MeSO <sub>2</sub> -dPhe-Pro-Arg-pNA)
		Pefachrome fVIIa (CH <sub>3</sub> SO <sub>2</sub> -D-CHA-But-Arg-pNA)

pNA = para-nitranilide (chromogenic)

35

30 AMC = amino methyl coumarin (fluorescent)

If none of the above substrates are cleaved, a coupled assay, described above, can be used. Briefly, test the ability of the protease to activate and enzyme, such as plasminogen and trypsinogen. To perform these assays, the single chain protease is incubated with a zymogen, such as plasminogen or trypsinogen, in the presence of the a known substrate, such, lys-plasminogen,

-162-

for the zymogen. If the single chain activates the zymogen, the activated enzyme, such as plasmin and trypsin, will degrade the substrate therefor.

#### **EXAMPLE 6**

#### Isolation and Cloning of Matriptase

5

10

15

20

25

30

#### A. Cell type and growth of cells

Human prostate adenocarcinoma cell line, PC-3, was purchased from ATCC (catalog number CRL-1435; Manassas, VA). The cells were cultured at 37°C, 5% CO<sub>2</sub> in Ham's F-12K growth medium (catalog number 9077; Irvine) supplemented with 2 mM L-glutamine and 10% fetal bovine serum. All subsequent cell manipulations were carried out according to the manufacturer's instructions. PC-3 cells were allowed to grow to about 90% confluence, and were then washed briefly with 1x phosphate buffered saline.

# B. Isolation of total RNA, and purification and enrichment of polyA<sup>+</sup> RNA

PC-3 cells were lysed in Trizol reagent (catalog number 15596; Life Technologies, Rockville, MD) and total RNA was isolated according to the manufacturer's protocol. The concentration of total RNA was estimated from absorbance reading at 260 nm. PolyA<sup>+</sup> RNA was purified and enriched using oligo-dT beads (catalog number 70061; Oligotex, Qiagen, Valencia, CA).

#### C. Reverse-transcription and polymerase chain reaction (PCR)

PC-3-derived polyA<sup>+</sup> RNA was converted to single-stranded cDNA (sscDNA) by reverse transcription using ProSTAR first-strand RT-PCR kit (catalog number 200420; Stratagene, La Jolla, CA) and SuperScript II RNase H<sup>-</sup> reverse transcriptase (catalog number 18064-022; Life Technologies). After reverse transcription, an aliquot of PC-3 sscDNA (4  $\mu$ L) was subjected to PCR using 2 mM each of the sense and anti-sense degenerate oligonucleotide primers and *Taq* polymerase (catalog number 201203; Qiagen). Total reaction volume was 100  $\mu$ L. The sequence of the sense primer was 5'-

TGGRT(I)VT(I)WS(I)GC(I)RC(I)CAYTG-3' (SEQ ID No. 13) and that of the antisense was 5'(I)GG(I)CC(I)CC(I)SWRTC(I)CCYT(I)RCA(I)GHRTC-3' (SEQ ID No. 14), where R=A,G; V=G,A,C; W=A,T; S=G,C; Y=C,T; H=A,T,C. The primer sequences correspond to two highly conserved regions in all

-163-

chymotrypsin-like serine proteases and amplify PCR products ranging from approximately 400 to 500 base pairs.

#### D. Clone screening and sequencing

10

15

20

25

30

The PCR products were separated on a 2% agarose gel and purified using a gel extraction kit (catalog number 28706; QIAquick gel extraction kit; Qiagen). The purified DNA fragments were ligated into pCR2.1-TOPO (catalog number K4500-01; Invitrogen, Carlsbad, CA). After transformation into E. coli cells, plasmid DNA was isolated and analyzed by digestion with EcoRI restriction enzyme. Clones that had inserted nucleic acid were further characterized by sequencing using a fluorescent dye-based DNA sequencing method (catalog number 4303149; BigDye terminator cycle sequencing kit with AmpliTaq DNA polymerase; Perkin Elmer, Lincoln, CA). A total of 31 clones were sequenced and analyzed. All sequences were analyzed by a multiple nucleotide sequence alignment algorithm (blastn) (www.ncbi.nlm.nih.gov/blast) to identify identical or closely related DNA deposited in GenBank (NCBI, Bethesda, MD). Those that did not show significant homology were further analyzed using blastx, which compares the six-frame conceptual translation products of a nucleotide sequence (both strands) against a protein sequence database (SwissProt). Eight clones yielded identical cDNA fragments that encode MTSP1. MTSP1 was subsequently found to be identical to matriptase (GenBank accession number AF118224).

## E. Rapid amplification of cDNA ends (RACE) and gene-specific amplification of MTSP1

To obtain DNA encoding the complete protease domain of MTSP1, RACE and gene-specific amplification reactions were performed. A human prostate Marathon-Ready cDNA (catalog # 7418-1; Clontech) was used to isolate part of the cDNA encoding MTSP1. Marathon-Ready cDNA is prepared to contain a known hybridization sequence at the 5' and 3' ends of the sscDNA. The 3' region of MTSP1 cDNA was obtained by a 3'-RACE reaction using a gene specific primer, 5'-CACCCCTTCTTCAATGACTTCACCTTCG-3' (SEQ ID No. 55). The 5' end of the MTSP1 protease domain was obtained by gene-specific amplification reaction using two MTSP1-specific primers, 5'-

-164-

TACCTCTCCTACGACTCC-3' (SEQ ID No. 56) for the sense primer and 5'-GAGGTTCTCGCAGGTGGTCTGGTTG-3' (SEQ ID No. 57) for the antisense primer. The sequences for these two primers were obtained from the human SNC19 mRNA sequence. The 3'-RACE reaction and gene-specific PCR produced DNA fragments that were >1 kbp in size. These fragments were subcloned into pCR2.1-TOPO (Invitrogen, San Diego, CA). After transformation into *E. coli* cells, plasmid DNA was isolated and analyzed by digestion with *Eco*RI restriction enzyme. Clones that had inserts were characterized by Southern blot analysis (using the internal cDNA fragment as probe) and by DNA sequence analysis.

## 10 F. PCR amplification of cDNA encoding the protease domain of MTSP1

To obtain a cDNA fragment encoding the entire protease domain of MTSP1, an end-to-end PCR amplification using gene-specific primers was used. The two primers used were: 5'-

15 CTCGAGAAAAGAGTTGTTGGGGGCACGGATGCGGATGAG-3' (SEQ ID No. 58) for the 5' end and 5'-GCGGCCGCACTATACCCCAGTGTTCTCTTTGATCCA-3' (SEQ ID No. 36 for the 3' end. The 5' primer contained the sequence that encodes the start of the MTSP1 protease domain (VVGGTDADE) (SEQ. ID. NO. 10). The 3' primer contained the stop codon of MTSP1. A ~800-bp fragment was amplified, purified and subcloned into the *Pichia pastoris* expression vector, pPIC9K, resulting in pPIC9K-MTSP1.

## G. Gene expression profile of MTSP1 in normal tissues, cancer cells and cancer tissues

To obtain information regarding the tissue distribution and gene
expression level of MTSP1, the DNA insert from pPIC9K-MTSP1 was used to probe a blot containing RNA from 76 different human tissues (catalog number 7775-1; human multiple tissue expression (MTE) array; CLONTECH, Palo Alto, CA). Significant expression was observed in the colon (ascending, transverse and descending), rectum, trachea, esophagus and duodenum. Moderate
expression levels were observed in the jejunum, ileum, ilocecum, stomach, prostate, pituitary gland, appendix, kidney, lung, placenta, pancreas, thyroid gland, salivary gland, mammary gland, fetal kidney, and fetal lung. Lower

-165-

expression levels were seen in the spleen, thymus, peripheral blood leukocyte, lymph node, bone marrow, bladder, uterus, liver, adrenal gland, fetal heart, fetal liver, fetal spleen, and fetal thymus. A significant amount of the MTSP1 transcript was also detected in colorectal adenocarcinoma cell line (SW480), Burkitt's lymphoma cell line (Daudi), and leukemia cell line (HL-60). RT-PCR of the MTSP1 transcript in several human primary tumors xenografted in athymic nude mice was performed using gene-specific primers. A high level of MTSP1 transcript was detected in colon adenocarcinoma (CX-1) and pancreatic adenocarcinoma (GI-103). Moderate levels were observed in another colon adenocarcinoma (GI-112), ovarian carcinoma (GI-102), lung carcinoma (LX-1), and breast carcinoma (GI-101). Another lung carcinoma (GI-117) expressed a low level of the MTSP1 transcript. A similar RT-PCR was performed to detect the presence of the MTSP1 transcript in PC-3 and LNCaP cell lines. Both cell lines expressed significant amounts of MTSP1 transcript.

#### H. Sequence analysis

10

15

25

30

All derived DNA and protein sequences were analyzed using MacVector (version 6.5; Oxford Molecular Ltd., Madison, WI). The cDNA encoding the protease domain of MTSP1 is composed of 726 base pairs which translate into a 241-amino acid protein sequence (rMAP) (see SEQ ID No. 1, 2, 49 and 50).

20 EXAMPLE 7

Production of Recombinant Serine Protease Domain of Matriptase or MTSP1 (rMAP)

#### A. Fermentation

The production of multi-milligram amounts of rMAP was carried out by fermentation in a BioFlo 3000 fermentor (New Brunswick Scientific, NJ) equipped with a 3.3 L capacity bioreactor using a SMD1168/pPIC9K:MTSP1 Sac SC1 clone. ZA001 complex media (10 g/L yeast extract, 20 g/L peptone, 40 g/L glycerol, 5 g/L ammonium sulfate, 0.2 g/L calcium sulfate dihydrate, 2 g/L magnesium sulfate heptahydrate, 2 g/L potassium sulfate, 25 g/L sodium hexametaposphate, 4.35 ml/L PTM1) was inoculated with 100 ml of an overnight culture of the *P. pastoris* transformant. The culture was supplemented

-166-

with 50% glycerol by fed-batch phase and induced for 18-24 hours with methanol controlled at 0.025%.

5

10

30

## B. Purification of Recombinant Serine Protease Domain of Matriptase or MTSP1 (rMAP)

The rMAP was secreted into the culture medium, so the first step of the purification involved the removal of cells and cell debris by centrifugation at 5000 g for 30 minutes. The resulting supernatant was decanted, adjusted to pH 8.0 with 10 N NaOH, and filtered through a SartoBran 300 0.45  $\pm$  0.2  $\mu$ M capsule. This supernatant was concentrated to 1 L by ultrafiltration using a 10 kDa ultrafiltration cartridge (NC SRT UF system with AG/Technologies UFP-10-C-5A filter), and the buffer was exchanged by crossflow filtration into 50 mM tris-HCl, 50 mM NaCl, 0.05% tween-80, pH 8.0 (buffer A). The filtration unit was rinsed once with 1 L buffer A which was combined with the concentrate.

The concentrated rMAP-containing solution was passed over a 150 ml benzamidine column that had been equilibrated with buffer A, at a flow rate of 8 ml/min. The column was washed with 3 column volumes of 50 mM tris-HCl, 1.0 M NaCl, 0.05% tween-80, pH 8.0 (buffer B) and eluted with 3 column volumes of 50 mM tris-HCl, 1.0 M L-arginine, 0.05% tween-80, pH 8.0 (buffer C). Fractions containing rMAP were identified by activity assay and pooled. This pooled material was concentrated to 10 ml using a JumboSep concentrator (Pall Gelman) and a 10 kDa cutoff membrane. Once concentrated to 10 ml, the buffer was exchanged into 50 mM Na<sub>2</sub>HPO<sub>4</sub>, 125 mM NaCl, pH 5.5 (buffer D) and the volume adjusted to 5-10 ml. The retentate was removed and the concentrator washed with buffer D which was added to the concentrate. The total sample volume was adjusted 15 ml.

The partially purified rMAP was passed through a 5 ml Q-sepharose Fast Flow HiTrap column (Amersham-Pharmacia Biotech) pre-equilibrated with 15 ml of buffer D. The flow through was collected. The HiTrap column was washed with an additional 10 ml of buffer D. Both flow throughs were pooled, and the protein concentration was determined by measurement of  $OD_{280}$  (using an extinction coefficient of  $2.012 \text{ mg/}OD_{280}$ ). Purified rMAP was then

-167-

deglycosylated by the addition 0.1  $\mu$ l of Endoglycosidase H (ProZyme, 5 U/ml) per mg of protein and incubating overnight at 4°C with gentle swirling.

The conductivity of the deglycosylated pool was adjusted to 2.0-3.0 mS/cm with Nanopure H<sub>2</sub>O and the pH adjusted to 6.5 (-200-300 mL final volume). The rMAP was then further purified by anion exchange chromatography by loading directly onto a Pharmacia Akta Explorer system using a 7 mL Source 15Q anion exchange column (Amersham-Pharmacia Biotech). The protein was eluted in a buffer containing 50 mM HEPES, pH 6.5 with a 0-0.33 M NaCl gradient over 10 column volumes at a flow rate of 6 ml/min.

10 Fractions containing protein were pooled, and benzamidine was added to a final concentration of 10 mM. Protein purity was examined by SDS-PAGE and protein concentration determined by measurement of OD<sub>280</sub> and use of a theoretical extinction coefficient of 2.012 mg/OD<sub>280</sub>.

#### **EXAMPLE 8**

#### 15 Assays

20

25

30

## Amidolytic Assay for Determining Inhibition of Serine Protease Activity of Matriptase or MTSP1

The ability of test compounds to act as inhibitors of rMAP catalytic activity was assessed by determining the inhibitor-induced inhibition of amidolytic activity by the MAP, as measured by IC<sub>50</sub> values. The assay buffer was HBSA (10 mM Hepes, 150mM sodium chloride, pH 7.4, 0.1% bovine serum albumin). All reagents were from Sigma Chemical Co. (St. Louis, MO), unless otherwise indicated.

Two IC $_{50}$  assays (a) one at either 30-minutes or 60-minutes (a 30-minute or a 60-minute preincubation of test compound and enzyme) and (b) one at 0-minutes (no preincubation of test compound and enzyme) were conducted. For the IC $_{50}$  assay at either 30-minutes or 60-minutes, the following reagents were combined in appropriate wells of a Corning microtiter plate: 50 microliters of HBSA, 50 microliters of the test compound, diluted (covering a broad concentration range) in HBSA (or HBSA alone for uninhibited velocity measurement), and 50 microliters of the rMAP (Corvas International) diluted in buffer, yielding a final enzyme concentration of 250 pM as determined by active

-168-

site filtration. Following either a 30-minute or a 60-minute incubation at ambient temperature, the assay was initiated by the addition of 50 microliters of the substrate S-2765 (N- $\alpha$ -Benzyloxycarbonyl-D-arginyl-L-glycyl-L-arginine-p-nitroaniline dihydrochloride; DiaPharma Group, Inc.; Franklin, OH) to each well, yielding a final assay volume of 200 microliters and a final substrate concentration of 100  $\mu$ M (about 4-times  $K_m$ ). Before addition to the assay mixture, S-2765 was reconstituted in deionized water and diluted in HBSA. For the IC<sub>50</sub> assay at 0 minutes; the same reagents were combined: 50 microliters of HBSA, 50 microliters of the test compound, diluted (covering the identical concentration range) in HBSA (or HBSA alone for uninhibited velocity measurement), and 50 microliters of the substrate S-2765. The assay was initiated by the addition of 50 microliters of rMAP. The final concentrations of all components were identical in both IC<sub>50</sub> assays (at 30- or 60- and 0-minute).

10

15

20

25

30

The initial velocity of chromogenic substrate hydrolysis was measured in both assays by the change of absorbance at 405 nM using a Thermo Max® Kinetic Microplate Reader (Molecular Devices) over a 5 minute period, in which less than 5% of the added substrate was used. The concentration of added inhibitor, which caused a 50% decrease in the initial rate of hydrolysis was defined as the respective  $IC_{50}$  value in each of the two assays (30- or 60-minutes and 0-minute).

### In vitro enzyme assays for specificity determination

The ability of compounds to act as a selective inhibitor of matriptase activity was assessed by determining the concentration of test compound that inhibits the activity of matriptase by 50%, ( $IC_{50}$ ) as described in the above Example, and comparing  $IC_{50}$  value for matriptase to that determined for all or some of the following serine proteases: thrombin, recombinant tissue plasminogen activator (rt-PA), plasmin, activated protein C, chymotrypsin, factor Xa and trypsin.

The buffer used for all assays was HBSA (10 mM HEPES, pH 7.5, 150 mM sodium chloride, 0.1% bovine serum albumin). The assay for  $IC_{50}$  determinations was conducted by combining in appropriate wells of a Corning microtiter plate, 50 microliters of HBSA, 50 microliters of the test compound at

-169-

a specified concentration (covering a broad concentration range) diluted in HBSA (or HBSA alone for  $V_0$  (uninhibited velocity) measurement), and 50 microliters of the enzyme diluted in HBSA. Following a 30 minute incubation at ambient temperature, 50 microliters of the substrate at the concentrations specified below were added to the wells, yielding a final total volume of 200 microliters. The initial velocity of chromogenic substrate hydrolysis was measured by the change in absorbance at 405 nm using a Thermo Max® Kinetic Microplate Reader over a 5 minute period in which less than 5% of the added substrate was used. The concentration of added inhibitor which caused a 50% decrease in the initial rate of hydrolysis was defined as the  $IC_{50}$  value.

Thrombin (flla) Assay

10

15

20

25

30

Enzyme activity was determined using the chromogenic substrate, Pefachrome t-PA (CH<sub>3</sub>SO<sub>2</sub>-D-hexahydrotyrosine-glycyl-L-Arginine-p-nitroaniline, obtained from Pentapharm Ltd.). The substrate was reconstituted in deionized water prior to use. Purified human α-thrombin was obtained from Enzyme Research Laboratories, Inc. .The buffer used for all assays was HBSA (10 mM HEPES, pH 7.5, 150 mM sodium chloride, 0.1% bovine serum albumin).

IC<sub>50</sub> determinations were conducted where HBSA (50  $\mu$ L),  $\alpha$ -thrombin (50  $\mu$ l) (the final enzyme concentration is 0.5 nM) and inhibitor (50  $\mu$ l) (covering a broad concentration range), were combined in appropriate wells and incubated for 30 minutes at room temperature prior to the addition of substrate Pefachrome-t-PA (50  $\mu$ l) (the final substrate concentration is 250  $\mu$ M, about 5 times Km). The initial velocity of Pefachrome t-PA hydrolysis was measured by the change in absorbance at 405 nm using a Thermo Max® Kinetic Microplate Reader over a 5 minute period in which less than 5% of the added substrate was used. The concentration of added inhibitor which caused a 50% decrease in the initial rate of hydrolysis was defined as the IC<sub>50</sub> value.

Factor Xa

Factor Xa catalytic activity was determined using the chromogenic substrate S-2765 (N-benzyloxycarbonyl-D-arginine-L-glycine-L-arginine-p-nitro-aniline), obtained from DiaPharma Group (Franklin, OH). All substrates were reconstituted in deionized water prior to use. The final concentration of S-2765

-170-

was 250  $\mu$ M (about 5-times Km). Purified human Factor X was obtained from Enzyme Research Laboratories, Inc. (South Bend, IN) and Factor Xa (FXa) was activated and prepared from it as described [Bock, P.E., Craig, P.A., Olson, S.T., and Singh, P. *Arch. Biochem. Biophys.* 273:375-388 (1989)]. The enzyme was diluted into HBSA prior to assay in which the final concentration was 0.25 nM. Recombinant tissue plasminogen activator (rt-PA) Assay

rt-PA catalytic activity was determined using the substrate, Pefachrome t-PA (CH<sub>3</sub>SO<sub>2</sub>-D-hexahydrotyrosine-glycyl-L-arginine-p-nitroaniline, obtained from Pentapharm Ltd.). The substrate was made up in deionized water followed by dilution in HBSA prior to the assay in which the final concentration was 500 micromolar (about 3-times Km). Human rt-PA (Activase®) was obtained from Genentech Inc. The enzyme was reconstituted in deionized water and diluted into HBSA prior to the assay in which the final concentration was 1.0 nM.

#### Plasmin Assay

10

20

25

30

Plasmin catalytic activity was determined using the chromogenic substrate, S-2366 [L-pyroglutamyl-L-prolyl-L-arginine-p-nitroaniline hydrochloride], which was obtained from DiaPharma group. The substrate was made up in deionized water followed by dilution in HBSA prior to the assay in which the final concentration was 300 micromolar (about 2.5-times Km).

Purified human plasmin was obtained from Enzyme Research Laboratories, Inc.

The enzyme was diluted into HBSA prior to assay in which the final concentration was 1.0 nM.

#### Activated Protein C (aPC) Assay

aPC catalytic activity was determined using the chromogenic substrate, Pefachrome PC (delta-carbobenzloxy-D-lysine-L-prolyl-L-arginine-p-nitroaniline dihydrochloride), obtained from Pentapharm Ltd.). The substrate was made up in deionized water followed by dilution in HBSA prior to the assay in which the final concentration was 400 micromolar (about 3-times Km). Purified human aPC was obtained from Hematologic Technologies, Inc. The enzyme was diluted into HBSA prior to assay in which the final concentration was 1.0 nM.

-171-

#### Chymotrypsin Assay

Chymotrypsin catalytic activity was determined using the chromogenic substrate, S-2586 (methoxy-succinyl-L-arginine-L-prolyl-L-tyrosyl-p-nitroanilide), which was obtained from DiaPharma Group. The substrate was made up in deionized water followed by dilution in HBSA prior to the assay in which the final concentration was 100 micromolar (about 9-times Km). Purified (3X-crystallized; CDI) bovine pancreatic alpha-chymotrypsin was obtained from Worthington Biochemical Corp. The enzyme was reconstituted in deionized water and diluted into HBSA prior to assay in which the final concentration was 0.5 nM.

#### 10 Trypsin Assay

Trypsin catalytic activity was determined using the chromogenic substrate, S-2222 (benzoyl-L-isoleucine-L-glutamic acid-[gamma-methyl ester]-L-arginine-p-nitroanilide), which was obtained from DiaPharma Group. The substrate was made up in deionized water followed by dilution in HBSA prior to the assay in which the final concentration was 250 micromolar (about 4-times Km). Purified (3X-crystallized; TRL3) bovine pancreatic trypsin was obtained from Worthington Biochemical Corp. The enzyme was reconstituted in deionized water and diluted into HBSA prior to assay in which the final concentration was 0.5 nM.

20

15

Since modifications will be apparent to those of skill in this art, it is intended that this invention be limited only by the scope of the appended claims.

5

15

20

#### WHAT IS CLAIMED IS:

- 1. A substantially purified single chain polypeptide, comprising the protease domain of a type-II membrane-type serine protease (MTSP) or a catalytically active portion thereof, wherein:
- the MTSP portion of the protein consists essentially of the protease domain of the MTSP or a catalytically active portion thereof.
- 2. The substantially purified polypeptide of claim 1, wherein the MTSP is not expressed on endothelial cells.
- The substantially purified polypeptide of claim 1, wherein the
   MTSP is not expressed on normal endothelial cells *in vivo*.
  - 4. The substantially purified polypeptide of claim 1, wherein the MTSP is a human protein.
  - 5. The substantially purified polypeptide of claim 1 that consists essentially of the protease domain of an MTSP or a catalytically active portion of the protease domain.
  - 6. The substantially purified polypeptide of claim 1, wherein the expression and/or activity of the MTSP in tumor cells differs from its level of expression and/or activity in non-tumor cells.
  - 7. The substantially purified polypeptide of claim 1, wherein the MTSP is detectable in a body fluid at a level that differs from its level in body fluids in a subject not having a tumor.
    - 8. The substantially purified polypeptide of claim 1, wherein: the MTSP is present in a tumor; and
- a substrate or cofactor for the MTSP is expressed at levels that differ from a non-tumor cell in the same tissue.
  - 9. The substantially purified polypeptide of claim 1, wherein: the MTSP exhibits altered substrate specificity in the tumor compared to its specificity in a non-tumor cell in the same tissue.
- 10. The substantially purified polypeptide of claim 1, wherein the 30 MTSP has an N-terminus that comprises IVNG, ILGG, VGLL or ILGG.
  - 11. The substantially purified polypeptide of claim 1, wherein the MTSP is selected from among MTSP1, MTSP3, MTSP4 and MTSP6.

- 12. The substantially purified polypeptide of claim 1, wherein the protease domain comprises the sequence of amino acids set forth as amino acids 615-855 of SEQ ID No. 2, as amino acids 205-437 of SEQ ID NO. 4, as the amino acids in SEQ ID No. 6, or as amino acids 217-443 in SEQ ID No. 12.
- 13. The substantially purified polypeptide of claim 1 that has at least about 40%, 60%, 80%, 90% or 95% sequence identity with a protease domain that comprises the sequence of amino acids set forth as amino acids 615-855 of SEQ ID No. 2, as amino acids 205-437 of SEQ ID NO. 4, as the amino acids in SEQ ID No. 6, or as amino acids 217-443 in SEQ ID No. 12.
- 14. A polypeptide of claim 1, wherein the protease domain portion is encoded by a nucleic acid molecule that hybridizes under conditions of high stringency along its full length to a nucleic acid molecule comprising a sequence of nucleotides set forth in SEQ ID No: 1, 3, 5, 7, 9 or 11 or to a molecule that encodes the protein set forth in SEQ ID No: 2, 4, 6, 8, 10 or 12 or at least one domain thereof.
  - 15. A nucleic acid molecule, comprising a sequence of nucleotides that encodes the polypeptide of claim 1.
- 16. A mutein of the polypeptide of claim 1, wherein:up to about 90% of the amino acids are replaced with another amino20 acid;

and the resulting polypeptide is a single chain and has catalytic activity at least 10% of the unmutated polypeptide.

- 17. The mutein of claim 16, wherein up to about 95% of the amino acids are replaced.
- 18. The mutein of claim 16, wherein the resulting polypeptide is a single chain and has catalytic activity at least 50% of the unmutated polypeptide.

25

- 19. A mutein of the polypeptide of claim 1, wherein a free Cys in the protease domain is replaced with another amino acid, whereby the resulting polypeptide exhibits proteolytic activity.
- 20. A mutein of the polypeptide of claim 1, wherein a free Cys in the protease domain is replaced with a serine.

#### -174-

- 21. A vector comprising the nucleic acid molecule of claim 15.
- 22. The vector of claim 21 that is an expression vector.
- 23. The vector of claim 21 that includes a sequence of nucleotides that directs secretion of any protein encoded by a sequence of nucleotides operatively linked thereto.
  - 24. The vector of claim 21 that is a Pichia vector or an E. coli vector.
  - 25. A cell, comprising the vector of claim 21.
  - 26. The cell of claim 25 that is a prokaryotic cell.
  - 27. The cell of claim 25 that is a eukaryotic cell.
- 10 28. The cell of claim 25 that is selected from among a bacterial cell, a yeast cell, a plant cell, an insect cell and an animal cell.
  - 29. The cell of claim 25 that is a mammalian cell.
  - 30. A method for producing a polypeptide that contains a protease domain of an MTSP, comprising:
- culturing the cell of claim 25 under conditions whereby the encoded protein is expressed by the cell; and

recovering the expressed protein.

5

- 31. The method of claim 30, wherein the cell is a pichia cell and the protein is secreted into the culture medium.
- 20 32 An antisense nucleic acid molecule that comprises at least 14 contiguous nucleotides or modified nucleotides that are complementary to a contiguous sequence of nucleotides in the protease domain of an MTSP of claim 1; or
- comprises at least 16 contiguous nucleotides or modified nucleotides that are complementary to a contiguous sequence of nucleotides in the protease domain of an MTSP of claim 1.
  - 33. An antibody that specifically binds to the single chain form of a protease domain of the polypeptide of claim 1, or a fragment or derivative of the antibody containing a binding domain thereof, wherein the antibody is a polyclonal antibody or a monoclonal antibody.

#### -175-

- 34. The polypeptide of claim 1, wherein the MTSP is selected from among corin, MTSP1, enterpeptidase, human airway trypsin-like protease (HAT), MTSP1, TMPRSS2, and TMPRSS4.
  - 35. A conjugate, comprising:
- 5 a) a protein of claim 1, and

- b) a targeting agent linked to the protein directly or via a linker.
- 142. The conjugate of claim 35, wherein the targeting agent permits
  - i) affinity isolation or purification of the conjugate;
  - ii) attachment of the conjugate to a surface;
  - iii) detection of the conjugate; or
  - iv) targeted delivery to a selected tissue or cell.
- 36. A combination, comprising:
- a) an inhibitor of the catalytic activity of the polypeptide of claim 1;
   and
- b) another treatment or agent selected from anti-tumor and antiangiogenic treatments or agents.
  - 37. The combination of claim 36, wherein the inhibitor and the antitumor and/or anti-angiogenic agent are formulated in a single pharmaceutical composition or each is formulated in separate pharmaceutical compositions.
- 20 38. The combination of claim 36, wherein the inhibitor is selected from antibodies and antisense oligonucleotides.
  - 39. A solid support comprising two or more polypeptides of claim 1 linked thereto either directly or via a linker.
- 40. The support of claim 39, wherein the polypeptides comprise an 25 array.
  - 41. The support of claim 39, wherein the polypeptides comprise a plurality of different MTSP protease domains.
  - 42. A method for identifying compounds that modulate the protease activity of an MTSP, comprising:
- contacting a polypeptide of claim 1 with a substrate proteolytically cleaved by the MTSP, and, either simultaneously, before or after, adding a test compound or plurality thereof;

-176-

measuring the amount of substrate cleaved in the presence of the test compound; and

selecting compounds that change the amount cleaved compared to a control, whereby compounds that modulate the activity of the MTSP are identified.

- 43. The method of claim 42, wherein the test compounds are small molecules, peptides, peptidomimetics, natural products, antibodies or fragments thereof.
- 44. The method of claim 42, wherein a plurality of the test substances are screened simultaneously.
  - 45. The method of claim 42, wherein the change in the amount cleaved is assessed by comparing the amount cleaved in the presence of the test compound with the amount in the absence of the test compound.
- 46. The method of claim 42, wherein a plurality of the test substances are screened for simultaneously.
  - 47. The method of claim 42, wherein a plurality of the polypeptides are linked to a solid support, either directly or via a linker.
  - 48. The method of claim 42, wherein the polypeptides comprise an array.
- 20 49. The method of claim 42, wherein the polypeptides comprise a plurality of different MTSP proteases.

25

30

50. A method of identifying a compound that specifically binds to a single chain protease domain of an MTSP, comprising:

contacting a polypeptide of claim 1 with a test compound or plurality thereof under conditions conducive to binding thereof; and

identifying compounds that specifically bind to the MTSP single chain protease domain or compounds that inhibit binding of a compound known to bind to the MTPS single chain protease domain, wherein the known compound is contacted with the polypeptide before, simultaneously with or after the test compound.

51. The method of claims 50, wherein the polypeptide is linked either directly or indirectly via a linker to a solid support.

-177-

- 52. The method of claim 50, wherein the test compounds are small molecules, peptides, peptidomimetics, natural products, antibodies or fragments thereof.
- 53. The method of claim 50, wherein a plurality of the test substances are screened for simultaneously.
  - 54. The method of claim 51, wherein a plurality of the polypeptides are linked to a solid support.

10

15

of:

- 55. A substantially purified membrane-type serine protease 3 (MTSP3).
- 56. The MTSP3 of claim 43 that is selected from the group consisting
- a polypeptide encoded by the sequence of nucleotides set forth in SEQ ID No. 3;
- a polypeptide encoded by a sequence of nucleotides that hybridizes under conditions of high stringency to the sequence of nucleotides set forth in SEQ ID No. 3;
- a polypeptide that comprises the sequence of amino acids set forth as amino acids 205-437 of SEQ ID No. 4;
- a polypeptide that comprises a sequence of amino acids having at least about 90% sequence identity with the sequence of amino acids set forth in SEQ ID No. 4; and
  - a polypeptide encoded by a splice variant of the sequence of nucleotides set forth in SEQ ID No. 3.
  - 57. A nucleic acid molecule, comprising a sequence of nucleotides that encodes the polypeptide of claim 56.
- 25 58. A vector comprising the nucleic acid molecule of claim 57.
  - 59. A cell, comprising the vector of claim 58.
  - 60. The cell of claim 59 that expresses the MTSP3 on its surface.
  - 61. The cell of claim 59 that is a prokaryotic cell.
  - 62. The cells of claim 59 that is a eukaryotic cell.
- 30 63. The cell of claim 59 that is selected from among a bacterial cell, a yeast cell, a yeast cell, a plant cell, an insect cell and an animal cell.
  - 64. The cell of claim 59 that is a mammalian cell.

- of amino acids set forth in SEQ ID No. 4; or a sequence of amino acids encoded by a sequence of nucleotides that hybridizes under conditions of high stringency to the sequence of nucleotides set forth in SEQ ID No. 3; or a protein having at least about 90% sequence identity with the sequence of amino acids set forth in SEQ ID No. 4 and retaining protease activity.
  - 66. A method for producing an MTSP3, comprising:
    culturing the cell of claim 59 under conditions whereby the encoded
    protein is expressed by the cell; and

10 recovering the expressed protein.

acid molecule of claim 57.

15

- 67. An antisense nucleic acid molecule that comprises at least 14 contiguous nucleotides or modified nucleotides that are complementary to a contiguous sequence of nucleotides in the nucleic acid molecule of claim 57; or comprises at least 16 contiguous nucleotides or modified nucleotides that are complementary to a contiguous sequence of nucleotides in the in the nucleic
- 68. An antibody that specifically binds to the MTSP of claim 57, or a fragment or derivative of the antibody containing a binding domain thereof, wherein the antibody is a polyclonal antibody or a monoclonal antibody.
- 20 69. A method for treating or preventing a neoplastic disease, in a mammal, comprising administering to a mammal an effective amount of an inhibitor of an MTSP3 of claim 55.
  - 70. The method of claim 69, wherein the inhibitor is an antibody that specifically binds to the MTSP3, or a fragment or derivative of the antibody containing a binding domain thereof, wherein the antibody is a polyclonal antibody or a monoclonal antibody.
    - 71. A substantially purified membrane-type serine protease 4 (MTSP4).
  - 72. The substantially purified MTSP4 that is an MTSP4-L or an MTSP4-S.
- 73. The MTSP4 of claim 71 that is selected from the group consistingof:

-179-

a polypeptide encoded by the sequence of nucleotides set forth in SEQ ID No. 7 or 9;

a polypeptide encoded by a sequence of nucleotides that hybridizes under conditions of high stringency to the sequence of nucleotides set forth in SEQ ID No. 7 or 9;

a polypeptide that comprises the sequence of amino acids set forth in SEQ ID No. 6, 8 or 10; and

a polypeptide encoded by a splice variant of the sequence of nucleotides set forth in SEQ ID No. 7 or 9.

10

15

25

5

- 74. The MTSP4 of claim 73 that is an MTSP4-L or an MTSP4-S.
- 75. A nucleic acid molecule, comprising a sequence of nucleotides that encodes the polypeptide of claim 73.
  - 76. A vector comprising the nucleic acid molecule of claim 74.
- 77. A cell, comprising the vector of claim 76.
  - 78. The cell of claim 77 that expresses the MTSP4 on its surface.
  - 79. The cell of claim 77 that is a prokaryotic cell.
  - 80. The cells of claim 77 that is a eukaryotic cell.
- 81. The cell of claim 77 that is selected from among a bacterial cell, a yeast cell, a yeast cell, a plant cell, an insect cell and an animal cell.
  - 82. The cell of claim 77 that is a mammalian cell.
  - 83. The cell of claim 77, wherein the MTSP4 comprises the sequence of amino acids set forth in SEQ ID No. 6, 8 or 10; or a sequence of amino acids encoded by a sequence of nucleotides that hybridizes under conditions of high stringency to the sequence of nucleotides set forth in SEQ ID No. 7 or 9; or a protein having at least about 90% sequence identity with the sequence of amino acids set forth in SEQ ID No. 8 or 10 and retaining protease activity.
- 84. A method for producing an MTSP4, comprising: culturing the cell of claim 77 under conditions whereby the encoded30 protein is expressed by the cell; and

recovering the expressed protein.

-180-

- 85. An antisense nucleic acid molecule that comprises at least 14 contiguous nucleotides or modified nucleotides that are complementary to a contiguous sequence of nucleotides in the nucleic acid molecule of claim 75 or
- comprises at least 16 contiguous nucleotides or modified nucleotides that are complementary to a contiguous sequence of nucleotides in the nucleic acid molecule of claim 75.

5

10

15

- 86. An antibody that specifically binds to the MTSP of claim 72, or a fragment or derivative of the antibody containing a binding domain thereof, wherein the antibody is a polyclonal antibody or a monoclonal antibody.
- 87. An antibody that specifically binds to the MTSP of claim 73, or a fragment or derivative of the antibody containing a binding domain thereof, wherein the antibody is a polyclonal antibody or a monoclonal antibody.
- 88. A method for treating or preventing a neoplastic disease, in a mammal, comprising administering to a mammal an effective amount of an inhibitor of an MTSP4 of claim 71.
- 89. The method of claim 88, wherein the inhibitor is an antibody that specifically binds to an MTSP4, or a fragment or derivative of the antibody containing a binding domain thereof, wherein the antibody is a polyclonal antibody or a monoclonal antibody.
- 20 90. A substantially purified membrane-type serine protease 6 (MTSP6) selected from the group consisting of:
  - a polypeptide encoded by the sequence of nucleotides set forth in SEQ ID No. 11;
  - a polypeptide encoded by a sequence of nucleotides that hybridizes along the full length thereof under conditions of high stringency to the sequence of nucleotides set forth in SEQ ID No. 11;
    - a polypeptide that comprises the sequence of amino acids set forth as amino acids 217-443 of SEQ ID No. 12;
- a polypeptide encoded by a splice variant of the sequence of nucleotides set forth in SEQ ID No. 11.
  - 91. A nucleic acid molecule, comprising a sequence of nucleotides that encodes the polypeptide of claim 90.

PCT/US01/03471

- 92. A vector comprising the nucleic acid molecule of claim 91.
- 93. A cell, comprising the vector of claim 92.
- 94. The cell of claim 93 that expresses the MTSP6 on its surface.
- 95. The cell of claim 93 that is a prokaryotic cell.
- 5 96. The cells of claim 93 that is a eukaryotic cell.
  - 97. The cell of claim 93 that is selected from among a bacterial cell, a yeast cell, a yeast cell, a plant cell, an insect cell and an animal cell.
    - 98. The cell of claim 93 that is a mammalian cell.
- 99. The cell of claim 93, wherein the MTSP6 comprises the sequence of amino acids set forth in SEQ ID No. 12; or a sequence of amino acids encoded by a sequence of nucleotides that hybridizes along the full length thereof under conditions of high stringency to the sequence of nucleotides set forth in SEQ ID No. 11; or a protein having at least about 95% sequence identity with the sequence of amino acids set forth in SEQ ID No. 12 and retaining protease activity.
  - 100. A method for producing an MTSP6, comprising:

culturing the cell of claim 93 under conditions whereby the encoded protein is expressed by the cell; and

recovering the expressed protein.

20 101. An antisense nucleic acid molecule that comprises at least 14 contiguous nucleotides or modified nucleotides that are complementary to a contiguous sequence of nucleotides in the nucleic acid molecule of claim 91; or

comprises at least 16 contiguous nucleotides or modified nucleotides that are complementary to a contiguous sequence of nucleotides in the in the nucleic acid molecule of claim 91.

- 102. An antibody that specifically binds to the MTSP of claim 90, or a fragment or derivative of the antibody containing a binding domain thereof, wherein the antibody is a polyclonal antibody or a monoclonal antibody.
- 103. A method for treating or preventing a neoplastic disease, in a mammal, comprising administering to a mammal an effective amount of an inhibitor of an MTSP6 of claim 90.

-182-

- 104. The method of claim 103, wherein the inhibitor is an antibody that specifically binds to the MTSP6, or a fragment or derivative of the antibody containing a binding domain thereof, wherein the antibody is a polyclonal antibody or a monoclonal antibody.
- 5 105. A recombinant non-human animal, wherein an endogenous gene of an MTSP has been deleted or inactivated by homologous recombination or insertional mutagenesis of the animal or an ancestor thereof.
  - 106. A recombinant non-human animal of claim 105, wherein the MTSP is an MTSP1, MTSP3, MTSP4 or MTSP6.
- 10 107. A conjugate, comprising:

15

20

25

30

- a) an MTSP3 or MTSP4 or an MTSP6 of claim 90; and
- b) a targeting agent linked to the protein directly or via a linker.
- 108. The conjugate of claim 106, wherein the targeting agent permits
  - i) affinity isolation or purification of the conjugate;
  - ii) attachment of the conjugate to a surface;
  - iii) detection of the conjugate; or
  - iv) targeted delivery to a selected tissue or cell.
- 109. A combination, comprising:
- a) an inhibitor of the catalytic activity of an MTSP3 or MTSP4 or MTSP6 of claim 90; and
- b) another treatment or agent selected from anti-tumor and antiangiogenic treatments or agents.
- 110. The combination of claim 109, wherein the inhibitor and the antitumor and/or anti-angiogenic agent are formulated in a single pharmaceutical composition or each is formulated in separate pharmaceutical compositions.

O

- 111. The combination of claim 109, wherein the inhibitor is selected from antibodies and antisense oligonucleotides.
- 112. A solid support comprising two or more MTSP3 polypeptides and/or MTSP4 polypeptides and/or MTSP6 polypeptides of claim 90 linked thereto either directly or via a linker.
- 113. The support of claim 112, wherein the polypeptides comprise an array.

114. A method for identifying compounds that modulate the protease activity of an MTSP selected from MTSP3, MTSP4 or MTSP6 of claim 90, comprising:

contacting the MTSP with a substrate proteolytically cleaved by the
 MTSP, and, either simultaneously, before or after, adding a test compound or plurality thereof;

measuring the amount of substrate cleaved in the presence of the test compound; and

selecting compounds that change the amount cleaved compared to a control, whereby compounds that modulate the activity of the MTSP are identified.

10

25

- 115. The method of claim 114, wherein the test compounds are small molecules, peptides, peptidomimetics, natural products, antibodies or fragments thereof.
- 15 117. The method of claim 114, wherein the change in the amount cleaved is assessed by comparing the amount cleaved in the presence of the test compound with the amount in the absence of the test compound.
  - 118. The method of claim 114, wherein a plurality of the test substances are screened for simultaneously.
- 20 119. The method of claim 118, wherein a plurality of the polypeptides are linked to a solid support.
  - 120. A method of identifying a compound that specifically binds to a MTSP selected from MTSP3, MTSP4 and the MTSP6 of claim 90, comprising: contacting the MTSP polypeptide with a test compound or plurality thereof under conditions conducive to binding thereof; and identifying compounds that specifically binds to the MTSP.
  - 121. A method of identifying a compound that specifically binds to a MTSP selected from MTSP3, MTSP4 and the MTSP6 of claim 90, comprising: contacting the MTSP polypeptide with a test compound or plurality thereof under conditions conducive to binding thereof; and

identifying compounds that specifically binds to the MTSP.

- 122. The method of any claims 121, wherein the polypeptide is linked either directly or indirectly via a linker to a solid support.
- 123. The method of claim 121, wherein the test compounds are small molecules, peptides, peptidomimetics, natural products, antibodies or fragments thereof.
  - 124. The method of claim 121, wherein a plurality of the test substances are screened for simultaneously.

5

15

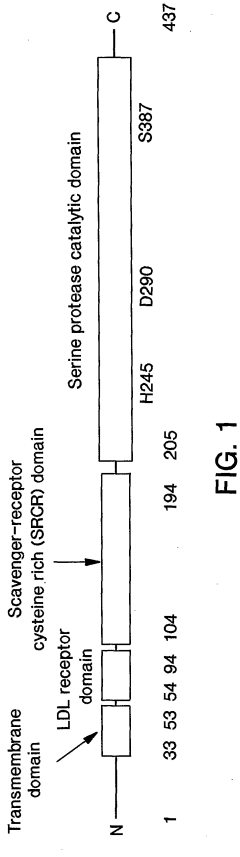
- 125. The method of claim 124, wherein a plurality of the polypeptides.10 are linked to a solid support.
  - 126. An MTSP6 polypeptide, comprising amino acids set forth as amino acids 46-55 in SEQ ID No. 12 and/or amino acids 368-394 of SEQ ID No. 12, and that is encoded by a sequence of nucleic acids that hybridizes under moderate stringency to nucleic acid encoding the polypeptide set forth in SEQ ID No. 12.
  - 127. The polypeptide of claim 126 that comprises the amino acids set forth as amino acids 46-55 in SEQ ID No. 12 and/or amino acids 368-394 of SEQ ID No. 12, and that is encoded by a sequence of nucleic acids that hybridizes under high stringency along its full length or full length of the protease domain to nucleic acid encoding the polypeptide set forth in SEQ ID No. 12.
  - 128. The polypeptide of claim 126, comprising the sequence of amino acids set forth in SEQ ID No. 12.
  - 129. An isolated nucleic acid molecule, comprising a sequence of nucleic acids that encodes the polypeptide of claim 126.
- 130. A method for treating tumors, comprising administering a prodrug that is specifically cleaved by an MTSP, whereby, upon contact with a cell that expresses MTSP activity, the prodrug is converted into an active drug.
  - 131. The method of claim 130, wherein the MTSP is selected from among an MTSP3, MTSP4 and an MTSP6 of claim 90.
- 30 132. The method of claim 130, wherein the MTSP is selected from among corin, enterpeptidase, human airway trypsin-like protease (HAT), MTSP1, TMPRSS2, and TMPRSS4.

-185-

- 133. A method of detecting neoplastic disease, comprising: detecting an MTSP3, MTSP4 or MTSP6 of claim 90 in a biological sample, wherein the amount detected differs from the amount in a subject who does not have neoplastic disease.
- 134. The method of claim 133, wherein the biological sample is selected from the group consisting of blood, urine, saliva, tears, interstitial fluid, cerebrospinal fluid, ascites fluid, tumor tissue biopsy and circulating tumor cells.
- 135. The method of claim 133, wherein the extracellular domain of the MTSP3, MTSP4 or MTSP6 is in the sample.
- 136. A modulator of the activity of a MTSP, which modulator is identified by the method of any of claims 42 and 114.

- 137. Use of the polyeptide of claim 1 for identifying compounds that modulate the activity of an MTSP.
- 138. The use of claim 137, wherein the compounds inhibit the15 proteolytic acitivity thereof.
  - 139 The use of claim 137, wherein the MTSP is selected from among MTSP1, MTSP3, MTSP4, MTSP6, corin, enterpeptidase, human airway trypsin-like protease (HAT), TMPRSS2, and TMPRSS4.
- 140. Use of a polypeptide of any of claims 55, 71 and 90 for treatment of a neoplastic disease.
  - 141. Use of a polypeptide of claim 55, 71 and 90 for formulation of a medicament of treatment of neoplastic disease.

Domain organization of MTSP3

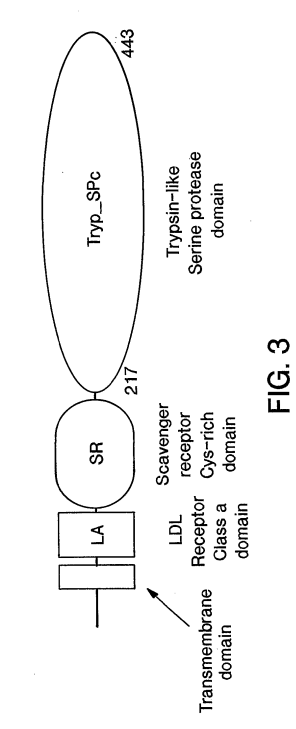


SUBSTITUTE SHEET (RULE 26)

S

S 802 658 Serine protease domain 448 481 521 567 304 337 377 423 LDLR domains Splice variant and domain organization of MTSP4 CUB domain 279 326 182 44 64 Transmembrane Z 136 domain 44 64 Z MTSP4-L (predominant) MTSP4-S **SUBSTITUTE SHEET (RULE 26)** 

Domain organization of human MTSP6



		$\nabla$		
MTSP3	194	LACGKSLKTPRVVGC	GEEASVDSWPWQVSIQYDKQHVCGGSILD	236
MTSP4-S	396	PQCDGRPDCRDGSDEEHCECGLQGPSSRIVGC	GAVSSEGEWPWQASLQVRGRHICGGALIA	455
MTSP4-L	540	PQCDGRPDCRDGSDEEHCECGLQGPSSRIVGC	GAVSSEGEWPWQASLQVRGRHICGGALIA	599
MTSP6	205	TACGHRRGYSSRIVGC 217		248
MTSP3	237	PHWVLTAAHCFRKHTDVFNWKVRAGSDKLO	GSFPSLAVAKIIIIEFNPMYPKDND	290
MTSP4-S	456	DRWVITAAHCFQEDSMASTVLWTVFLGKVWQN	NSRWPGEVSFKVSRLLLHPYHEEDSHDYD	515
MTSP4-L	600	DRWVITAAHCFQEDSMASTVLWTVFLGKVWQN	NSRWPGEVSFKVSRLLLHPYHEEDSHDYD	718
MTSP6	249	PLWIITAAHCVYDLYLPKSWTIQVGLVSLI	LDNPAPSHLVEKIVYHSKYKPKRLGND	304
MTSP3	291	IALMKLQFPLTFSGTVRPICLPFFDEELTPAT	rplwiigwgftkQnggkmsdillQasvQv	350
MTSP4-S	516	VALLQLDHPVVRSAAVRPVCLPARSHFFEPGI	LHCWITGWGALRE-GGPISNALQKVDVQL	574
MTSP4-L	660	VALLQLDHPVVRSAAVRPVCLPARSHFFEPGI	LHCWITGWGALRE-GGPISNALQKVDVQL	718
MTSP6	305	IALMKLAGPLTFNEMIQPVCLPNSEENFPDGF	CVCWTSGWGATED-GGDASPVLNHAAVPL	363
MTSP3	351	IDSTRCNADDAYQGEVTEKMMCAGIPEGGVDT	CCQGDSGGPLMYQSDQWHVVGIVSWGY	408
MTSP4-S	575	IPQDLCSEVYRYQVTPRMLCAGYRKGKKD	ACQGDSGGPLVCKALSGRWFLAGLVSWGL	632
MTSP4-L	719	IPQDLCSEVYRYQVTPRMLCAGYRKGKKDA	ACQGDSGGPLVCKALSGRWFLAGLVSWGL	776
MTSP6	364	ISNKICNHRDVYGGIISPSMLCAGYLTGGVDS	GCQGDSGPLVCQERR-LWKVLVGATSFGI	442
MTSP3	409	GCGGPSTPGVYTKVSAYLNWIYNVWKAEL	437	
MTSP4-S	633	GCGRPNYFGVYTRITGVISWIQQVVT	658	
MTSP4-L ·	777	GCGRPNYFGVYTRITGVISWIQQVVT	802	
MTSP6	423	GCAEVNKPGVYTRVTSFLDWIHEQMERDLKT	453	

## √ cleavage site

- + potential glycosylation site
- \* unpaired cysteine

## FIG. 4

## SEQUENCE LISTING

<110> Edwin L. Madison Edgar O. Ong Jiunn-Chern Yeh Corvas International, Inc. <120> NUCLEIC ACID MOLECULES ENCODING TRANSMEMBRANE SERINE PROTEASES, THE ENCODED PROTEINS AND METHODS BASED THEREON <130> 24745-1607 <140> 09/000,000 <141> 2001-02-01 <150> 60/213,124 <151> 2000-06-22 <150> 60/234,840 <151> 2000-06-22 <150> 60/179,982 <151> 2000-02-03 <150> 60/183,542 <151> 2000-02-18 <150> 09/657,968 <151> 2000-02-08 <160> 72 <170> FastSEQ for Windows Version 4.0 <210> 1 <211> 3147 <212> DNA <213> Homo Sapien <220> <223> Nucleotide encoding MTSP1 <221> CDS <222> (23)...(2589) <301> O'Brien, T.J. and Tanimoto, H. <308> GenBank AR081724 <310> US Pat 5972616 <311> 1998-02-20 <312> 1999-10-26 tcaagagegg ceteggggta ee atg ggg age gat egg gee ege aag gge gga 52 Met Gly Ser Asp Arg Ala Arg Lys Gly Gly ggg ggc ccg aag gac ttc ggc gcg gga ctc aag tac aac tcc cgg cac 100 Gly Gly Pro Lys Asp Phe Gly Ala Gly Leu Lys Tyr Asn Ser Arg His

gag Glu	aaa Lys	gtg Val	aat Asn 30	ggc Gly	ttg Leu	gag Glu	gaa Glu	ggc Gly 35	gtg Val	gag Glu	ttc Phe	ctg Leu	cca Pro 40	gtc Val	aac Asn	148
	gtc Val															196
	gcc Ala 60															244
ctg Leu 75	gtg Val	tgg Trp	cat His	ttg Leu	cag Gln 80	tac Tyr	cgg Arg	gac Asp	gtg Val	cgt Arg 85	gtc Val	cag Gln	aag Lys	gtc Val	ttc Phe 90	292
	ggc Gly															340
	tcc Ser															388
	ctg Leu															436
aag Lys	gag Glu 140	tcg Ser	gct Ala	gtg Val	acg Thr	gcc Ala 145	ttc Phe	agc Ser	gag Glu	ggc Gly	agc Ser 150	gtc Val	atc Ile	gcc Ala	tac Tyr	484
	tgg Trp															532
	gtc Val															580
tcc Ser	ctg Leu	aag Lys	tcc Ser 190	ttt Phe	gtg Val	gtc Val	acc Thr	tca Ser 195	gtg Val	gtg Val	gct Ala	ttc Phe	ccc Pro 200	acg Thr	gac Asp	628
	aaa Lys		Val			Thr		Asp					Phe			676
cac His	gcc Ala 220	cgc Arg	ggt Gly	gtg Val	gag Glu	ctg Leu 225	atg Met	cgc Arg	ttc Phe	acc Thr	acg Thr 230	ccc Pro	ggc Gly	ttc Phe	cct Pro	724
gac Asp 235	agc Ser	ccc Pro	tac Tyr	ccc Pro	gct Ala 240	cat His	gcc Ala	cgc Arg	tgc Cys	cag Gln 245	tgg Trp	gcc Ala	ctg Leu	cgg Arg	999 Gly 250	772
	gcc Ala															820
tcc Ser	tgc Cys	gac Asp	gag Glu	cgc Arg	ggc Gly	agc Ser	gac Asp	ctg Leu	gtg Val	acg Thr	gtg Val	tac Tyr	aac Asn	acc Thr	ctg Leu	868

	270			275			280		
	gag Glu								916
	aac Asn								964
	acc Thr								1012
	ctg Leu								1060
	aca Thr 350								1108
	tgc Cys								1156
	ttc Phe								1204
	ccc Pro								1252
	tcc Ser								1300
	cac His 430								1348
	tcc Ser								1396
	Gly ggg								1444
	acc Thr								1492
	ttc Phe								1540
	agt Ser 510								1588

					cag Gln											163	36
					tgc Cys											168	34
gac Asp 555	gag Glu	gcc Ala	tcc Ser	tgc Cys	ccc Pro 560	aag Lys	gtg Val	aac Asn	gtc Val	gtc Val 565	act Thr	tgt Cys	acc Thr	aaa Lys	cac His 570	173	32
					aat Asn											178	30
tgt Cys	gac Asp	gly aaa	aag Lys 590	gag Glu	gac Asp	tgt Cys	agc Ser	gac Asp 595	ggc Gly	tca Ser	gat Asp	gag Glu	aag Lys 600	gac Asp	tgc Cys	182	28
gac Asp	tgt Cys	605 Gly 999	ctg Leu	cgg Arg	tca Ser	ttc Phe	acg Thr 610	aga Arg	cag Gln	gct Ala	cgt Arg	gtt Val 615	gtt Val	Gl <sup>A</sup> aaa	ggc Gly	187	76
acg Thr	gat Asp 620	gcg Ala	gat Asp	gag Glu	ggc Gly	gag Glu 625	tgg Trp	ccc Pro	tgg Trp	cag Gln	gta Val 630	agc Ser	ctg Leu	cat His	gct Ala	192	24
					atc Ile 640											197	72
					cac His											202	20
					tgg Trp											206	58
					gl <sup>à</sup> aaa											211	16
					aat Asn											216	54
					ccg Pro 720											221	L2
					tcc Ser											226	50
gtc Val	acg Thr	Gly ggc	tgg Trp 750	gga Gly	cac His	acc Thr	cag Gln	tat Tyr 755	gga Gly	ggc Gly	act Thr	ggc Gly	gcg Ala 760	ctg Leu	atc Ile	230	80
ctg Leu	caa Gln	aag Lys	ggt Gly	gag Glu	atc Ile	cgc Arg	gtc Val	atc Ile	aac Asn	cag Gln	acc Thr	acc Thr	tgc Cys	gag Glu	aac Asn	235	56

		765					770					775				
														ttc Phe		2404
agc Ser 795	ggc Gly	Gly ggc	gtg Val	gac Asp	tcc Ser 800	tgc Cys	cag Gln	ggt Gly	gat Asp	tcc Ser 805	gly aaa	gga Gly	ccc Pro	ctg Leu	tcc Ser 810	2452
agc Ser	gtg Val	gag Glu	gcg Ala	gat Asp 815	gly aaa	cgg Arg	atc Ile	ttc Phe	cag Gln 820	gcc Ala	ggt Gly	gtg Val	gtg Val	agc Ser 825	tgg Trp	2500
														agg Arg		2548
							aaa Lys 850						ta g	3 <b>3</b> 330	cgggg	2599
ggct ctcc ggga agac cccc ggct agga cttc aaaa <210 <211 <212	cagages aggs aggs acages acage	gac to get a	ggad ccaa agggg cccc caagg cctgg	ceget atete gagga cegec gaget gaget cagac cttgt	eg ac ge et ac ac ca go ag eg gt go	etgea agaa etggt eccea gggaa gggc gtet	accas aaacc eggtt aagct acgga cctts gagas	g ego c tet c eta c ego a ego ego a ego ego ego ego ego ego ego ego ego ego	cccc ccgct actga gccga ctcga ccacq aaatt	aga tcc iccc iccc iggc jagc jagc jctc	acat tcag aact gcgt ctcc ttga	acad gcctd gggg ttgt tcag ggaa ccag	etg tegen	egaac aagtg aaagg atatc aaggt ccagg	ectgca etcaat ggagct gtttga etgcct eggtgg gctcgg ggtgga aaaaaa	2659 2719 2779 2839 2899 2959 3079 3139 3147
<400	)> 2		_													
Met 1	Gly	Ser	Asp	Arg 5	Ala	Arg	Lys	Gly	Gly 10	Gly	Gly	Pro	Lys	Asp 15	Phe	
Gly	Ala	Gly	Leu 20	Lys	Tyr	Asn	Ser	Arg 25	His	Glu	Lys	Val	Asn 30	Gly	Leu	
Glu	Glu	Gly 35		Glu	Phe	Leu	Pro 40		Asn	Asn	Val	Lys 45		Val	Glu	
Lys	His		Pro	Gly	Arg	Trp 55		Val	Leu		Ala 60		Leu	Ile	Gly	
Leu 65	Leu	Leu	Val	Leu	Leu 70		Ile	${\tt Gl}_{\it Y}$	Phe			Trp	His	Leu		
	Arg	Asp	Val			Gln	Lys	Val			Gly	Tyr	Met	Arg	80 Ile	
Thr	Asn	Glu		85 Phe	Val	Asp	Ala		90 Glu	Asn	Ser	Asn		95 Thr	Glu	
Phe	Val	Ser	100 Leu	Ala	Ser	Lys		105 Lys	Asp	Ala	Leu		110 Leu	Leu	Tyr	
Ser		115 Val	Pro	Phe	Leu		120 Pro	Tyr	His	Lys		125 Ser	Ala	Val	Thr	
Ala	130	Val			Ser	135	Pro			Tyr	140	Ser		Val Phe	Ser	
Ala 145	130 Phe	Val Ser	Glu	Gly Leu	Ser 150	135 Val	Pro Ile	Ala	Tyr Glu	Tyr 155	140 Trp	Ser Ser	Glu	Phe Glu	Ser 160	
Ala 145 Ile	130 Phe Pro	Val Ser Gln	Glu His	Gly Leu 165	Ser 150 Val	135 Val Glu	Pro Ile Glu	Ala Ala	Tyr Glu 170	Tyr 155 Arg	140 Trp Val	Ser Ser Met	Glu Ala	Phe	Ser 160 Glu	

Val	Thr	Ser	180 Val	Val	Ala	Phe	Pro	185 Thr	Asp	Ser	Lys	Thr	190 Val	Gln	Arg
		195	Asn				200					205			
	210		Phe			215					220				
225			Cys		230					235					240
			Phe	245					250					255	
			260 Val					265	,				270		
	_	275	Gln				280					285			
	290					295					300				
305			Ser		310					315					320
			His	325					330					335	
Met	Ser	Ser	Cys 340	Gly	Gly	Arg	Leu	Arg 345	Lys	Ala	Gln	Gly	Thr 350	Phe	Asn
Ser	Pro	Tyr 355	Tyr	Pro	Gly	His	Tyr 360	Pro	Pro	Asn	Ile	Asp 365	Cys	Thr	Trp
Asn	Ile 370	Glu	Val	Pro	Asn	Asn 375	Gln	His	Val	Lys	Val 380	Ser	Phe	Lys	Phe
Phe 385	Tyr	Leu	Leu	Glu	Pro 390	Gly	Val	Pro	Ala	Gly 395	Thr	Cys	Pro	Lys	Asp 400
Tyr	Val	Glu	Ile	Asn 405	Gly	Glu	Lys	Tyr	Cys 410	Gly	Glu	Arg	Ser	Gln 415	Phe
Val	Val	Thr	Ser 420	Asn	Ser	Asn	Lys	Ile 425	Thr	Val	Arg	Phe	His 430	Ser	Asp
Gln	Ser	Tyr 435	Thr	Asp	Thr	Gly	Phe 440	Leu	Ala	Glu	Tyr	Leu 445	Ser	Tyr	Asp
Ser	Ser 450	Asp	Pro	Cys	Pro	Gly 455	Gln	Phe	Thr	Cys	Arg 460	Thr	Gly	Arg	Cys
Ile 465	Arg	Lys	Glu	Leu	Arg 470	Cys	Asp	Gly	Trp	Ala 475	Asp	Cys	Thr	Asp	His 480
Ser	Asp	Glu	Leu	Asn 485	Cys	Ser	Cys	Asp	Ala 490	Gly	His	Gln	Phe	Thr 495	Cys
Lys	Asn	Lys	Phe 500	Cys	Lys	Pro	Leu	Phe 505	Trp	Val	Cys	Asp	Ser 510	Val	Asn
Asp	Cys	Gly 515	Asp	Asn	Ser	Asp	Glu 520	Gln	Gly	Cys	Ser	Cys 525	Pro	Ala	Gln
Thr	Phe 530	Arg	Cys	Ser	Asn	Gly 535	Lys	Cys	Leu	Ser	Lys 540	Ser	Gln	Gln	Cys
Asn 545	Gly	Lys	Asp	Asp	Cys 550	Gly	Asp	Gly	Ser	Asp 555	Glu	Ala	Ser	Cys	Pro 560
	Val	Asn	Val	Val 565		Cys	Thr	Lys	His 570		Tyr	Arg	Cys	Leu 575	
Gly	Leu	Сув	Leu 580		Tys	Gly	Asn	Pro 585		Cys	Asp	Gly	Lys 590		Asp
Cys	Ser	Asp 595	Gly	Ser	Asp	Glu	Lys 600		Cys	Asp	Cys	Gly 605		Arg	Ser
Phe	Thr 610		Gln	Ala	Arg	Val 615		Gly	Gly	Thr	Asp 620		Asp	Glu	Gly
Glu 625		Pro	Trp	Gln	Val 630		Leu	His	Ala	Leu 635		Gln	Gly	His	Ile 640
	Gly	Ala	Ser	Leu 645		Ser	Pro	Asn	Trp 650		Val	Ser	Ala	Ala 655	
Cys	Tyr	Ile	Asp 660		Arg	Gly	Phe	Arg 665		Ser	Asp	Pro	Thr 670		Trp
Thr	Ala	Phe	Leu	Gly	Leu	His	Asp		Ser	Gln	Arg	Ser		Pro	Gly

```
675
                              680
Val Gln Glu Arg Arg Leu Lys Arg Ile Ile Ser His Pro Phe Phe Asn
    690
                          695
                                                700
Asp Phe Thr Phe Asp Tyr Asp Ile Ala Leu Leu Glu Leu Glu Lys Pro
                      710
                                           715
Ala Glu Tyr Ser Ser Met Val Arg Pro Ile Cys Leu Pro Asp Ala Ser
                 725
                                       730
                                                             735
His Val Phe Pro Ala Gly Lys Ala Ile Trp Val Thr Gly Trp Gly His
             740
                                   745
                                                        750
Thr Gln Tyr Gly Gly Thr Gly Ala Leu Ile Leu Gln Lys Gly Glu Ile
        755
                              760
                                                    765
Arg Val Ile Asn Gln Thr Thr Cys Glu Asn Leu Leu Pro Gln Gln Ile
                          775
                                                780
Thr Pro Arg Met Met Cys Val Gly Phe Leu Ser Gly Gly Val Asp Ser
                      790
                                           795
Cys Gln Gly Asp Ser Gly Gly Pro Leu Ser Ser Val Glu Ala Asp Gly
                 805
                                       810
                                                             815
Arg Ile Phe Gln Ala Gly Val Val Ser Trp Gly Asp Gly Cys Ala Gln
                                   825
                                                        830
Arg Asn Lys Pro Gly Val Tyr Thr Arg Leu Pro Leu Phe Arg Asp Trp
        835
                              840
Ile Lys Glu Asn Thr Gly Val
    850
                          855
<210> 3
<211> 2137
<212> DNA
<213> Homo Sapien
<220>
<221> CDS
<222> (261)...(1574)
<223> DNA sequence encoding a transmembrane serine
     protease (MTSP3) protein
<400> 3
ccatectaat acgaeteact atagggeteg ageggeegee egggeaggte agagagagge
                                                                            60
agcagettge teageggaca aggatgetgg gegtgaggga ceaaggeetg eeetgeacte
                                                                           120
gggcctcctc cagccagtgc tgaccaggga cttctgacct gctggccagc caggacctgt
gtggggaggc cctcctgctg ccttggggtg acaatctcag ctccaggcta cagggagacc
                                                                           240
gggaggatca cagagccagc atg tta cag gat cct gac agt gat caa cct ctg
                                                                           293
                        Met Leu Gln Asp Pro Asp Ser Asp Gln Pro Leu
aac agc ctc gat gtc aaa ccc ctg cgc aaa ccc cgt atc ccc atg gag
                                                                           341
Asn Ser Leu Asp Val Lys Pro Leu Arg Lys Pro Arg Ile Pro Met Glu
acc ttc aga aag gtg ggg atc ccc atc atc ata gca cta ctg agc ctg Thr Phe Arg Lys Val Gly Ile Pro Ile Ile Ile Ala Leu Leu Ser Leu
                                                                           389
                               35
gcg agt atc atc att gtg gtt gtc ctc atc aag gtg att ctg gat aaa
                                                                           437
Ala Ser Ile Ile Ile Val Val Leu Ile Lys Val Ile Leu Asp Lys
tac tac ttc ctc tgc ggg cag cct ctc cac ttc atc ccg agg aag cag
                                                                           485
Tyr Tyr Phe Leu Cys Gly Gln Pro Leu His Phe Ile Pro Arg Lys Gln
                       65
ctg tgt gac gga gag ctg gac tgt ccc ttg ggg gag gac gag gag cac
Leu Cys Asp Gly Glu Leu Asp Cys Pro Leu Gly Glu Asp Glu Glu His
                                                                           533
```

80 85 90 581 tgt gtc aag agc ttc ccc gaa ggg cct gca gtg gca gtc cgc ctc tcc Cys Val Lys Ser Phe Pro Glu Gly Pro Ala Val Ala Val Arg Leu Ser aag gac cga tcc aca ctg cag gtg ctg gac tcg gcc aca ggg aac tgg Lys Asp Arg Ser Thr Leu Gln Val Leu Asp Ser Ala Thr Gly Asn Trp 629 11.5 tte tet qee tqt tte qae aac tte aca gaa get ete get gag aca gee 677 Phe Ser Ala Cys Phe Asp Asn Phe Thr Glu Ala Leu Ala Glu Thr Ala 130 tgt agg cag atg ggc tac agc agc aaa ccc acc ttc aga gct gtg gag 725 Cys Arg Gln Met Gly Tyr Ser Ser Lys Pro Thr Phe Arg Ala Val Glu 773 att ggc cca gac cag gat ctg gat gtt gtt gaa atc aca gaa aac agc Ile Gly Pro Asp Gln Asp Leu Asp Val Val Glu Ile Thr Glu Asn Ser 165 cag gag ctt cgc atg cgg aac tca agt ggg ccc tgt ctc tca ggc tcc Gln Glu Leu Arg Met Arg Asn Ser Ser Gly Pro Cys Leu Ser Gly Ser 821 869 ctg gtc tcc ctg cac tgt ctt gcc tgt ggg aag agc ctg aag acc ccc Leu Val Ser Leu His Cys Leu Ala Cys Gly Lys Ser Leu Lys Thr Pro 195 917 cgt gtg gtg ggt ggg gag gac tct gtg gat tct tgg cct tgg cag Arg Val Val Gly Gly Glu Glu Ala Ser Val Asp Ser Trp Pro Trp Gln 205 210 215 965 gtc agc atc cag tac gac ata cag cac gtc tgt gga ggg agc atc ctg Val Ser Ile Gln Tyr Asp Ile Gln His Val Cys Gly Gly Ser Ile Leu 225 235 gac ecc cac tgg gtc etc acg gea gee cac tgc ttc agg aaa cat acc 1013 Asp Pro His Trp Val Leu Thr Ala Ala His Cys Phe Arg Lys His Thr gat gtg ttc aac tgg aag gtg cgg gca ggc tca gac aaa ctg ggc agc 1061 Asp Val Phe Asn Trp Lys Val Arg Ala Gly Ser Asp Lys Leu Gly Ser 1109 ttc cca tcc ctg gct gtg gcc aag atc atc atc att gaa ttc aac ccc Phe Pro Ser Leu Ala Val Ala Lys Ile Ile Ile Glu Phe Asn Pro 1157 atg tac ccc aaa gac aat gac atc gcc ctc atg aag ctg cag ttc cca Met Tyr Pro Lys Asp Asn Asp Ile Ala Leu Met Lys Leu Gln Phe Pro 290 ctc act ttc tca ggc aca gtc agg ctc atc tgt ctg ccc ttc ttt gat 1205 Leu Thr Phe Ser Gly Thr Val Arg Leu Ile Cys Leu Pro Phe Phe Asp 305 315 gaq gaq ctc act cca gcc acc cca ctc tqq atc att gga tgg ggc ttt 1253

Glu Glu Leu Thr Pro Ala Thr Pro Leu Trp Ile Ile Gly Trp Gly Phe

acg aag cag aat gga ggg aag atg tct gac ata ctg ctg cag gcg tca Thr Lys Gln Asn Gly Gly Lys Met Ser Asp Ile Leu Leu Gln Ala Ser 335 340 345	1301
gtc cag gtc att gac agc aca cgg tgc aat gca gac gat gcg tac cag Val Gln Val Ile Asp Ser Thr Arg Cys Asn Ala Asp Asp Ala Tyr Gln 350 355 360	1349
ggg gaa gtc acc gag aag atg atg tgt gca ggc atc ccg gaa ggg ggt Gly Glu Val Thr Glu Lys Met Met Cys Ala Gly Ile Pro Glu Gly Gly 365 370 375	1397
gtg gac acc tgc cag ggt gac agt ggt ggg ccc ctg atg tac caa tct Val Asp Thr Cys Gln Gly Asp Ser Gly Gly Pro Leu Met Tyr Gln Ser 380 385 390 395	1445
gac cag tgg cat gtg gtg ggc atc gtt agc tgg ggc tat ggc tgc ggg Asp Gln Trp His Val Val Gly Ile Val Ser Trp Gly Tyr Gly Cys Gly 400 405 410	1493
ggc ccg agc acc cca gga gta tac acc aag gtc tca gcc tat ctc aac Gly Pro Ser Thr Pro Gly Val Tyr Thr Lys Val Ser Ala Tyr Leu Asn 415 420 425	1541
tgg atc tac aat gtc tgg aag gct gag ctg taa tgctgctgcc cctttgcagt Trp Ile Tyr Asn Val Trp Lys Ala Glu Leu * 430 435	1594
getgggagee getteettee tgeeetgeee acetggggat ceceaaagt cagacacaga geaagagtee cettgggtae acecetetge cacaageete ageetteett ggageageaa aggeeteaa tteetgtaag agaceetege ageecagagg cgeeagagg aagteageag cectageteg gecacaettg gtgeteecag cateecaggg agagacacag eccaetgaac aggetgtettg taaaageeca gateactgtg ggetggagag gagaaggaaa gggtetgege cageeetgte egtetteace cateeceaag cetactagag caagaaacea gttgtaatat aaaatgeact geeetactgt tggtatgaet acegttaeet actgttgtea ttgttattae agetatggee actattatta aagagetgtg taacaaaaaa aaaaaaaaaa aaaaaaaaaa aaaaaaaa	1654 1714 1774 1834 1894 1954 2014 2074 2134 2137
<210> 4 <211> 437 <212> PRT <213> Homo Sapien	
<pre>&lt;400&gt; 4 Met Leu Gln Asp Pro Asp Ser Asp Gln Pro Leu Asn Ser Leu Asp Val 1 5 10 15</pre>	
Lys Pro Leu Arg Lys Pro Arg Ile Pro Met Glu Thr Phe Arg Lys Val 20 25 30	
Gly Ile Pro Ile Ile Ile Ala Leu Leu Ser Leu Ala Ser Ile Ile Ile . 35 40 45	
Val Val Leu Ile Lys Val Ile Leu Asp Lys Tyr Tyr Phe Leu Cys 50 55 60	
Gly Gln Pro Leu His Phe Ile Pro Arg Lys Gln Leu Cys Asp Gly Glu 65 70 75 80	
Leu Asp Cys Pro Leu Gly Glu Asp Glu Glu His Cys Val Lys Ser Phe 85 90 95	
Pro Glu Gly Pro Ala Val Ala Val Arg Leu Ser Lys Asp Arg Ser Thr 100 105 110	
Leu Gln Val Leu Asp Ser Ala Thr Gly Asn Trp Phe Ser Ala Cys Phe 115 120 125	
Asp Asn Phe Thr Glu Ala Leu Ala Glu Thr Ala Cys Arg Gln Met Gly	

```
135
Tyr Ser Ser Lys Pro Thr Phe Arg Ala Val Glu Ile Gly Pro Asp Gln
145 150 155 160
Asp Leu Asp Val Val Glu Ile Thr Glu Asn Ser Gln Glu Leu Arg Met
                165
                                    170
                                                         175
Arg Asn Ser Ser Gly Pro Cys Leu Ser Gly Ser Leu Val Ser Leu His
           180
                                185
Cys Leu Ala Cys Gly Lys Ser Leu Lys Thr Pro Arg Val Val Gly Gly
                            200
Glu Glu Ala Ser Val Asp Ser Trp Pro Trp Gln Val Ser Ile Gln Tyr
                                          220
                        215
Asp Ile Gln His Val Cys Gly Gly Ser Ile Leu Asp Pro His Trp Val
                    230
                                        235
Leu Thr Ala Ala His Cys Phe Arg Lys His Thr Asp Val Phe Asn Trp
                245
                                    250
Lys Val Arg Ala Gly Ser Asp Lys Leu Gly Ser Phe Pro Ser Leu Ala
           260
                               265
Val Ala Lys Ile Ile Ile Glu Phe Asn Pro Met Tyr Pro Lys Asp
                            280
                                                 285
Asn Asp Ile Ala Leu Met Lys Leu Gln Phe Pro Leu Thr Phe Ser Gly
                        295
                                           300
Thr Val Arg Leu Ile Cys Leu Pro Phe Phe Asp Glu Glu Leu Thr Pro
                   310
                                        315
Ala Thr Pro Leu Trp Ile Ile Gly Trp Gly Phe Thr Lys Gln Asn Gly
                325
                                    330
                                                        335
Gly Lys Met Ser Asp Ile Leu Leu Gln Ala Ser Val Gln Val Ile Asp
                             345
            340
                                                   350
Ser Thr Arg Cys Asn Ala Asp Asp Ala Tyr Gln Gly Glu Val Thr Glu
        355
                            360
                                                 365
Lys Met Met Cys Ala Gly Ile Pro Glu Gly Gly Val Asp Thr Cys Gln
                        375
                                            380
Gly Asp Ser Gly Gly Pro Leu Met Tyr Gln Ser Asp Gln Trp His Val
                    390
                                         395
Val Gly Ile Val Ser Trp Gly Tyr Gly Cys Gly Gly Pro Ser Thr Pro
               405
                                  410
Gly Val Tyr Thr Lys Val Ser Ala Tyr Leu Asn Trp Ile Tyr Asn Val
           420
                                425
Trp Lys Ala Glu Leu
       435
<210> 5
<211> 708
<212> DNA
<213> Homo Sapien
<220>
<221> CDS
<222> (1)...(708)
<223> MTSP4 protease domain cDNA
<400> 5
att gtt ggt gga gct gtg tcc tcc gag ggt gag tgg cca tgg cag gcc Ile Val Gly Gly Ala Val Ser Ser Glu Gly Glu Trp Pro Trp Gln Ala
                                                                        96
age etc eag gtt egg ggt ega eac atc tgt ggg ggg gee etc atc get
Ser Leu Gln Val Arg Gly Arg His Ile Cys Gly Gly Ala Leu Ile Ala
gac cgc tgg gtg ata aca gct gcc cac tgc ttc cag gag gac agc atg
                                                                       144
Asp Arg Trp Val Ile Thr Ala Ala His Cys Phe Gln Glu Asp Ser Met
```

			gtg Val													192
			cct Pro													240
			cac His													288
			cac His 100													336
			cgc Arg													384
			ggc Gly													432
			gat Asp													480
			cag Gln													528
ggc	aag Lys	aag Lys	gat Asp 180	gcc Ala	tgt Cys	cag Gln	ggt Gly	gac Asp 185	tca Ser	ggt Gly	ggt Gly	ccg Pro	ctg Leu 190	gtg Val	tgc Cys	576
			agt Ser													624
			ggc ggc													672
	٧al	Ile	agc Ser	Trp	Ile	Gln					tga *					708
<212	-> 23 2> PF	$\mathbf{T}$	Sapie	∋n												
<400		<b>G</b> 7	a -			~	_	<b>~</b> 7	<b>~</b> ?	<b>~</b> 7	-	~	_	<b>~</b> 7	<b>73.7</b> -	
1			Gly	5					10		_			15		
Ser	Leu	Gln	Val 20	Arg	Gly	Arg	His	Ile 25	Cys	Gly	Gly	Ala	Leu 30	Ile	Ala	
Asp	Arg	Trp	Val	Ile	Thr	Ala	Ala	His	Cys	Phe	Gln	Glu	Asp	Ser	Met	

Ala																
	Ser 50	Thr	Val	Leu	Trp	Thr 55	Val	Phe	Leu	Gly	Lys 60	Val	Trp	Gln	Asn	
Ser 65	Arg	Trp	Pro	Gly	Glu 70	Val	Ser	Phe	Lys	Val 75	Ser	Arg	Leu	Leu	Leu 80	
	Pro	Tyr	His	Glu 85	Glu	Asp	Ser	His	Asp 90	Tyr	Asp	Val	Ala	Leu 95	Leu	
Gln	Leu	Asp	His 100	Pro	Val	Val	Arg	Ser 105	Ala	Ala	Val	Arg	Pro 110	Val	Cys	
Leu	Pro	Ala 115	Arg	Ser	His	Phe	Phe 120	Glu	Pro	Gly	Leu	His 125	Cys	Trp	Ile	
Thr	Gly 130	Trp	Gly	Ala	Leu	Arg 135	Glu	Gly	Gly	Pro	Ile 140	Ser	Asn	Ala	Leu	
Gln 145	Lys	Val	Asp	Val	Gln 150	Leu	Ile	Pro	Gln	Asp 155	Leu	Cys	Ser	Glu	Val 160	
Tyr	Arg	Tyr	Gln	Val 165	Thr	Pro	Arg	Met	Leu 170	Cys	Ala	Gly	Tyr	Arg 175	Lys	
Gly	Lys	Lys	Asp 180	Ala	Cys	Gln	Gly	Asp 185	Ser	Gly	Gly	Pro	Leu 190	Val	Cys	
Lys	Ala	Leu 195	Ser	Gly	Arg	Trp	Phe 200	Leu	Ala	Gly	Leu	Val 205	Ser	Trp	Gly	
Leu	Gly 210	Сув	Gly	Arg	Pro	Asn 215	Tyr	Phe	Gly	Val	Tyr 220	Thr	Arg	Ile	Thr	
Gly 225	Val	Ile	Ser	Trp	Ile 230	Gln	Gln	Val	Val	Thr 235						
<213 <213	0> 7 1> 3: 2> DI 3> Ho	AV	Sapie	en												
<222	0> 1> CI 2> (: 3> cI	33).	(24 enco	441) ding	: MTS	SP4-1	ت (lo	ong i	form)	sp]	Lice	var:	iant			
	0> 7 tagge	rca (	radd	rtaat	-c ac	rtaac	rcaca	a acr	ata	ccc	ata	acc	asa	acc	ccc	53
cca	99	JUA :	34995	gega	ع مر	gegas	goage	a ag			Val					35
cag Gln	gtg Val	gct Ala 10	ggc Gly	ggg Glv	cag	āāa	gac	aas								
		7.0		2	GTII	GIY	Asp 15	Gly	ggt Gly	gat Asp	ggc Gly	gag Glu 20	gaa Glu	gcg Ala	gag Glu	101
	gag Glu 25	<b>aaa</b>		ttc	aag	gcc	Asp 15 tgt	Gly	Gly gac	Asp	Gly	Glu 20 aga	Glu	Ala	-Glu cgg	101
Pro	Glu	ggg Gly ctc	Met	ttc Phe ctg	aag Lys gtg	gcc Ala 30	Asp 15 tgt Cys	gag Glu ttt	gac Asp gtg	Asp tcc Ser	aag Lys 35	Glu 20 aga Arg gcc	Glu aaa Lys ctg	Ala gcc Ala ctc	cgg Arg	
ggc Gly 40	Glu 25 tac	ggg Gly ctc Leu	Met cgc Arg	ttc Phe ctg Leu	aag Lys gtg Val 45	gcc Ala 30 ccc Pro	Asp 15 tgt Cys ctg Leu	gag Glu ttt Phe	gac Asp gtg Val	Asp tcc Ser ctg Leu 50	aag Lys 35 ctg Leu	glu 20 aga Arg gcc Ala	aaa Lys ctg Leu	gcc Ala ctc Leu	cgg Arg gtg Val 55	149
ggc Gly 40 ctg Leu	Glu 25 tac Tyr	ggg Gly ctc Leu tcg Ser	Met cgc Arg gcg Ala gtc	ttc Phe ctg Leu ggg Gly 60	aag Lys gtg Val 45 gtg Val	gcc Ala 30 ccc Pro cta Leu	Asp 15 tgt Cys ctg Leu ctc Leu	gag Glu ttt Phe tgg Trp	gac Asp gtg Val tat Tyr 65	tcc ser ctg Leu 50 ttc Phe agt	aag Lys 35 ctg Leu cta	gcc Ala ggg gly	aaa Lys ctg Leu tac Tyr	gcc Ala ctc Leu aag Lys 70 ctc	cgg Arg gtg Val 55 gcg Ala	149 197

		90					95					100				
			gcc Ala					Met								389
acc Thr 120	cgc Arg	ctg Leu	gga Gly	act Thr	tac Tyr 125	tac Tyr	aac Asn	tcc Ser	agc Ser	tcc Ser 130	gtc Val	tat Tyr	tcc Ser	ttt Phe	999 Gly 135	437
			ctc Leu													485
			ctg Leu 155													533
			ctg Leu													581
gcc Ala	gag Glu 185	tac Tyr	gaa Glu	gtg Val	gac Asp	ccc Pro 190	gag Glu	ggc Gly	cta Leu	gtg Val	atc Ile 195	ctg Leu	gaa Glu	gcc Ala	agt Ser	629
			ata Ile													677
			ggc Gly													725
ctg Leu	gcc Ala	tcc Ser	agc Ser 235	tgc Cys	ctg Leu	tgg Trp	cac His	ctg Leu 240	cag Gln	ggc Gly	ccc Pro	aag Lys	gac Asp 245	ctc Leu	atg Met	773
ctc Leu	aaa Lys	ctc Leu 250	cgg Arg	ctg Leu	gag Glu	tgg Trp	acg Thr 255	ctg Leu	gca Ala	gag Glu	tgc Cys	cgg Arg 260	gac Asp	cga Arg	ctg Leu	821
			gac Asp													869
gtg Val 280	tac Tyr	ggc Gly	tgc Cys	agc Ser	cgc Arg 285	cag Gln	gag Glu	ccc Pro	gtg Val	gtg Val 290	gag Glu	gtt Val	ctg Leu	gcg Ala	tcg Ser 295	917
			atg Met													965
			gtg Val 315													1013
			acg Thr													1061

					agc Ser											1109
					tct Ser 365											1157
gcc Ala	tat Tyr	gca Ala	ctg Leu	agg Arg 380	agg Arg	cag Gln	aag Lys	tat Tyr	gat Asp 385	ttg Leu	ccg Pro	tgc Cys	acc Thr	cag Gln 390	ggc Gly	1205
					aac Asn											1253
					atc Ile											1301
					atc Ile											1349
					cag Gln 445											1397
					tgt Cys											1445
					gag Glu											1493
					aca Thr											1541
cag Gln	cct Pro 505	gat Asp	tgt Cys	ctc Leu	aac Asn	ggc Gly 510	agc Ser	gac Asp	gaa Glu	gag Glu	cag Gln 515	tgc Cys	cag Gln	gaa Glu	gly aaa	1589
					ttc Phe 525											1637
aag Lys	aag Lys	ccc Pro	aac Asn	ccg Pro 540	cag Gln	tgt Cys	gat Asp	gly aaa	cgg Arg 545	ccc Pro	gac Asp	tgc Cys	agg Arg	gac Asp 550	ggc Gly	1685
					tgt Cys											1733
					gtg Val											1781
agc Ser	ctc Leu	cag Gln	gtt Val	cgg Arg	ggt Gly	cga Arg	cac His	atc Ile	tgt Cys	gly aaa	gly ggg	gcc Ala	ctc Leu	atc Ile	gct Ala	1829

585	590		595	
			cag gag gac agc atg Gln Glu Asp Ser Met 615	1877
			aag gtg tgg cag aac Lys Val Trp Gln Asn 630	1925
tcg cgc tgg cct Ser Arg Trp Pro 635	gga gag gtg tc Gly Glu Val Se	c ttc aag gtg r Phe Lys Val 640	agc cgc ctg ctc ctg Ser Arg Leu Leu 645	1973
		r His Asp Tyr	gac gtg gcg ctg ctg Asp Val Ala Leu Leu 660	2021
			gtg cgc ccc gtc tgc Val Arg Pro Val Cys 675	2069
			ctg cac tgc tgg att Leu His Cys Trp Ile 695	2117
			atc agc aac gct ctg Ile Ser Asn Ala Leu 710	2165
cag aaa gtg gat Gln Lys Val Asp 715	gtg cag ttg at Val Gln Leu Il	c cca cag gac e Pro Gln Asp 720	ctg tgc agc gag gtc Leu Cys Ser Glu Val 725	2213
		g Met Leu Cys	gcc ggc tac cgc aag Ala Gly Tyr Arg Lys 740	2261
ggc aag aag gat Gly Lys Lys Asp 745	gcc tgt cag gg Ala Cys Gln Gl 750	t gac tca ggt y Asp Ser Gly	ggt ccg ctg gtg tgc Gly Pro Leu Val Cys 755	2309
aag gca ctc agt Lys Ala Leu Ser 760	ggc cgc tgg tt Gly Arg Trp Ph 765	c ctg gcg ggg e Leu Ala Gly 770	ctg gtc agc tgg ggc Leu Val Ser Trp Gly 775	2357
			tac acc cgc atc aca Tyr Thr Arg Ile Thr 790	2405
ggt gtg atc agc Gly Val Ile Ser 795			tga ggaactgccc *	2451
ggacaagtat tetge tettgttteg teed ggggteaaga egted eteteeteeg teed atgetggtte tacai tgggeageet tged	gegggg ggtggggggggggggggggggggggggggggg	ag agagcaggcc ta tggcaggagg gc ccacacccag gc ctaatgcaag tg aggtgcgcc tc cagcttcgga	agggcaactg ccaagcaggg ctgtggtggc aggaggggca atgagaagtg ccagcagttg ccettttgcc tcccaattct gcagtggctc agcagcaga cactctgtac agaggctgtt agcccctggt ctaacttggg gagccctgga gactgccagg	2511 2571 2631 2691 2751 2811 2871 2931

tgggcctgct gccactgtaa gccaaaaggt ggggaagtcc tgactccagg gtccttgccc cacccctgcc tgccacctgg gccctcacag cccagaccct cactgggagg tgagctcagc <210> 8 <211> 802 <212> PRT <213> Homo Sapien Met Pro Val Ala Glu Ala Pro Gln Val Ala Gly Gln Gly Asp Gly Gly Asp Gly Glu Glu Ala Glu Pro Glu Gly Met Phe Lys Ala Cys Glu Asp Ser Lys Arg Lys Ala Arg Gly Tyr Leu Arg Leu Val Pro Leu Phe Val Leu Leu Ala Leu Leu Val Leu Ala Ser Ala Gly Val Leu Leu Trp Tyr Phe Leu Gly Tyr Lys Ala Glu Val Met Val Ser Gln Val Tyr Ser Gly Ser Leu Arg Val Leu Asn Arg His Phe Ser Gln Asp Leu Thr Arg Arg Glu Ser Ser Ala Phe Arg Ser Glu Thr Ala Lys Ala Gln Lys Met Leu Lys Glu Leu Ile Thr Ser Thr Arg Leu Gly Thr Tyr Tyr Asn Ser Ser Ser Val Tyr Ser Phe Gly Glu Gly Pro Leu Thr Cys Phe Phe Trp Phe Ile Leu Gln Ile Pro Glu His Arg Arg Leu Met Leu Ser Pro Glu Val Val Gln Ala Leu Leu Val Glu Glu Leu Leu Ser Thr Val Asn Ser Ser Ala Ala Val Pro Tyr Arg Ala Glu Tyr Glu Val Asp Pro Glu Gly Leu Val Ile Leu Glu Ala Ser Val Lys Asp Ile Ala Ala Leu Asn Ser Thr Leu Gly Cys Tyr Arg Tyr Ser Tyr Val Gly Gln Gly Gln Val Leu Arg Leu Lys Gly Pro Asp His Leu Ala Ser Ser Cys Leu Trp His Leu Gln Gly Pro Lys Asp Leu Met Leu Lys Leu Arg Leu Glu Trp Thr Leu Ala Glu Cys Arg Asp Arg Leu Ala Met Tyr Asp Val Ala Gly Pro Leu Glu Lys Arg Leu Ile Thr Ser Val Tyr Gly Cys Ser Arg Gln Glu Pro Val Val Glu Val Leu Ala Ser Gly Ala Ile Met Ala Val Val Trp Lys Lys Gly Leu His Ser Tyr Tyr Asp Pro Phe Val Leu Ser Val Gln Pro Val Val Phe Gln Ala Cys Glu Val Asn Leu Thr Leu Asp Asn Arg Leu Asp Ser Gln Gly Val Leu Ser Thr Pro Tyr Phe Pro Ser Tyr Tyr Ser Pro Gln Thr His Cys Ser Trp His Leu Thr Val Pro Ser Leu Asp Tyr Gly Leu Ala Leu Trp Phe Asp Ala Tyr Ala Leu Arg Arg Gln Lys Tyr Asp Leu Pro Cys Thr Gln Gly Gln Trp Thr Ile Gln Asn Arg Arg Leu Cys Gly Leu Arg Ile Leu Gln Pro Tyr Ala Glu Arg Ile Pro Val Val

```
Ala Thr Ala Gly Ile Thr Ile Asn Phe Thr Ser Gln Ile Ser Leu Thr
          420
                             425
Gly Pro Gly Val Arg Val His Tyr Gly Leu Tyr Asn Gln Ser Asp Pro
       435
                          440
Cys Pro Gly Glu Phe Leu Cys Ser Val Asn Gly Leu Cys Val Pro Ala
                  455
Cys Asp Gly Val Lys Asp Cys Pro Asn Gly Leu Asp Glu Arg Asn Cys
                                     . 475
465
                 470
Val Cys Arg Ala Thr Phe Gln Cys Lys Glu Asp Ser Thr Cys Ile Ser
              485
                                 4.90
Leu Pro Lys Val Cys Asp Gly Gln Pro Asp Cys Leu Asn Gly Ser Asp
           500
                              505
                                                510
Glu Glu Gln Cys Gln Glu Gly Val Pro Cys Gly Thr Phe Thr Phe Gln
       515
                          520
                                             525
Cys Glu Asp Arg Ser Cys Val Lys Lys Pro Asn Pro Gln Cys Asp Gly
                     535
                                        540
Arg Pro Asp Cys Arg Asp Gly Ser Asp Glu Glu His Cys Glu Cys Gly
                 550
                                 555
Leu Gln Gly Pro Ser Ser Arg Ile Val Gly Gly Ala Val Ser Ser Glu
                               570
            565
Gly Glu Trp Pro Trp Gln Ala Ser Leu Gln Val Arg Gly Arg His Ile
           580
                              585
                                                 590
Cys Gly Gly Ala Leu Ile Ala Asp Arg Trp Val Ile Thr Ala Ala His
595 600 605
               600
       595
                                             605
Cys Phe Gln Glu Asp Ser Met Ala Ser Thr Val Leu Trp Thr Val Phe
          615
                                         620
Leu Gly Lys Val Trp Gln Asn Ser Arg Trp Pro Gly Glu Val Ser Phe
                  630
                                     635
Lys Val Ser Arg Leu Leu His Pro Tyr His Glu Glu Asp Ser His
               645
                                 650
Asp Tyr Asp Val Ala Leu Leu Gln Leu Asp His Pro Val Val Arg Ser
           660
                             665
                                                 670
Ala Ala Val Arg Pro Val Cys Leu Pro Ala Arg Ser His Phe Phe Glu
      675
                         680
                                           685
Pro Gly Leu His Cys Trp Ile Thr Gly Trp Gly Ala Leu Arg Glu Gly
  690
                     695
                                         700
Gly Pro Ile Ser Asn Ala Leu Gln Lys Val Asp Val Gln Leu Ile Pro
705
                  710
                                      715
Gln Asp Leu Cys Ser Glu Val Tyr Arg Tyr Gln Val Thr Pro Arg Met
                                  730
               725
                                                    735
Leu Cys Ala Gly Tyr Arg Lys Gly Lys Lys Asp Ala Cys Gln Gly Asp
           740
                              745
Ser Gly Gly Pro Leu Val Cys Lys Ala Leu Ser Gly Arg Trp Phe Leu 755 760 765
                          760
Ala Gly Leu Val Ser Trp Gly Leu Gly Cys Gly Arg Pro Asn Tyr Phe
                      775
                                         780
Gly Val Tyr Thr Arg Ile Thr Gly Val Ile Ser Trp Ile Gln Gln Val
                                      795
785
                   790
Val Thr
<210> 9
<211> 2672
<212> DNA
<213> Homo Sapien
<220>
<221> CDS
<222> (33)...(2009)
```

<223> cDNA encoding: MTSP4-S (short form) splice variant

<400> 9

tcat	cgg	cca g	gaggg	gtgat	cc ag	gtgag	gcaga	a ag						gcc Ala		53
cag Gln	gtg Val	gct Ala 10	ggc Gly	61 <sup>7</sup> 888	cag Gln	G1 <sup>A</sup> aaa	gac Asp 15	gga Gly	ggt Gly	gat Asp	ggc Gly	gag Glu 20	gaa Glu	gcg Ala	gag Glu	101
						gcc Ala 30										149
ggc Gly 40	tac Tyr	ctc Leu	cgc Arg	ctg Leu	gtg Val 45	ccc Pro	ctg Leu	ttt Phe	gtg Val	ctg Leu 50	ctg Leu	gcc Ala	ctg Leu	ctc Leu	gtg Val 55	197
						cta Leu										245
gag Glu	gtg Val	atg Met	gtc Val 75	agc Ser	cag Gln	gtg Val	tac Tyr	tca Ser 80	gly ggc	agt Ser	ctg Leu	cgt Arg	gta Val 85	ctc Leu	aat Asn	293
						ctt Leu										· 341
						cag Gln 110										389
						tac Tyr										437
gtg Val	tac Tyr	ggc Gly	tgc Cys	agc Ser 140	cgc Arg	cag Gln	gag Glu	ccc Pro	gtg Val 145	gtg Val	gag Glu	gtt Val	ctg Leu	gcg Ala 150	tcg Ser	485
glà aaa	gcc Ala	atc Ile	atg Met 155	gcg Ala	gtc Val	gtc Val	tgg Trp	aag Lys 160	aag Lys	ggc Gly	ctg Leu	cac His	agc Ser 165	tac Tyr	tac Tyr	533
gac Asp	ccc Pro	ttc Phe 170	Val	ctc Leu	tcc Ser	gtg Val	cag Gln 175	Pro	gtg Val	gtc Val	Phe	cag Gln 180	Ala	tgt Cys	gaa Glu	581
gtg Val	aac Asn 185	ctg Leu	acg Thr	ctg Leu	gac Asp	aac Asn 190	agg Arg	ctc Leu	gac Asp	tcc Ser	cag Gln 195	gl <sup>y</sup> ggc	gtc Val	ctc Leu	agc Ser	629
						tac Tyr										677
cac His	ctc Leu	acg Thr	gtg Val	ccc Pro 220	tct Ser	ctg Leu	gac Asp	tac Tyr	ggc Gly 225	ttg Leu	gcc Ala	ctc Leu	tgg Trp	ttt Phe 230	gat Asp	725
gcc Ala	tat Tyr	gca Ala	ctg Leu	agg Arg	agg Arg	cag Gln	aag Lys	tat Tyr	gat Asp	ttg Leu	ccg Pro	tgc Cys	acc Thr	cag Gln	ggc Gly	773

			235					240					245			
						agg Arg										821
ccc Pro	tac Tyr 265	gcc Ala	gag Glu	agg Arg	atc Ile	ccc Pro 270	gtg Val	gtg Val	gcc Ala	acg Thr	gcc Ala 275	gl <sup>à</sup> aaa	atc Ile	acc Thr	atc Ile	869
						tcc Ser										917
tat Tyr	gly ggc	ttg Leu	tac Tyr	aac Asn 300	cag Gln	tcg Ser	gac Asp	ccc Pro	tgc Cys 305	cct Pro	gga Gly	gag Glu	ttc Phe	ctc Leu 310	tgt Cys	965
						gtc Val										1013
ccc Pro	aac Asn	ggc Gly 330	ctg Leu	gat Asp	gag Glu	aga Arg	aac Asn 335	tgc Cys	gtt Val	tgc Cys	aga Arg	gcc Ala 340	aca Thr	ttc Phe	cag Gln	1061
						tgc Cys 350										1109
cag Gln 360	cct Pro	gat Asp	tgt Cys	ctc Leu	aac Asn 365	ggc Gly	agc Ser	gac Asp	gaa Glu	gag Glu 370	cag Gln	Çys Çys	cag Gln	gaa Glu	999 Gly 375	1157
gtg Val	cca Pro	tgt Cys	gjå aaa	aca Thr 380	ttc Phe	acc Thr	ttc Phe	cag Gln	tgt Cys 385	gag Glu	gac Asp	cgg Arg	agc Ser	tgc Cys 390	gtg Val	1205
						tgt Cys										1253
						gaa Glu										1301
att Ile	gtt Val 425	ggt Gly	gga Gly	gct Ala	gtg Val	tcc Ser 430	tcc Ser	gag Glu	ggt Gly	gag Glu	tgg Trp 435	cca Pro	tgg Trp	cag Gln	gcc Ala	1349
						cga Arg										1397
						gct Ala										1445
gcc Ala	tcc Ser	acg Thr	gtg Val 475	ctg Leu	tgg Trp	acc Thr	gtg Val	ttc Phe 480	ctg Leu	ggc Gly	aag Lys	gtg Val	tgg Trp 485	cag Gln	aac Asn	1493

			cct Pro													1541
			cac His													1589
cag Gln 520	ctc Leu	gac Asp	cac His	ccg Pro	gtg Val 525	gtg Val	cgc Arg	tcg Ser	gcc Ala	gcc Ala 530	gtg Val	cgc Arg	ccc Pro	gtc Val	tgc Cys 535	1637
			cgc Arg													1685
acg Thr	ggc gly	tgg Trp	ggc Gly 555	gcc Ala	ttg Leu	cgc Arg	gag Glu	ggc Gly 560	ggc Gly	ccc Pro	atc Ile	agc Ser	aac Asn 565	gct Ala	ctg Leu	1733
cag Gln	aaa Lys	gtg Val 570	gat Asp	gtg Val	cag Gln	ttg Leu	atc Ile 575	cca Pro	cag Gln	gac Asp	ctg Leu	tgc Cys 580	agc Ser	gag Glu	gtc Val	1781
tat Tyr	cgc Arg 585	tac Tyr	cag Gln	gtg Val	acg Thr	cca Pro 590	cgc Arg	atg Met	ctg Leu	tgt Cys	gcc Ala 595	ggc Gly	tac Tyr	cgc Arg	aag Lys	1829
			gat Asp													1877
			agt Ser													1925
			ggc Gly 635													1973
			agc Ser								tga *	ggaa	actgo	ccc		2019
ggad toti gggg ctci atgg atci tggg cac	caagt cgttt gtcaa ccctq gcago cggga gcctq	tat i	tetge tecet tecee tacat tgeet ggaag	geggg cgatg cttc cccq ccaq ggtgq ctgta	gg gg gt ct ga gg ct co ga ga ga ga ct co aa go	gtggg cgtc gacce gagtg agcag ccate ccate	gggag cagta cagga gctga gtcta gatta cggag caca caca	g aga a tg c cc c ct g aga c ca g gg t gg g cc	agcag gcagg acacg actg gtgc gctt gacc ggaag caga	ggcc gagg ccag caag gccc cgga ctca gtcc ccct	ctgi atga ccci gcaci agco agco tgaco caci	tggtgaag ttttgg tetgg tetgg ceetg ceetg	ggc aggc aggc aggg aggg	aggag ccag tcccag agcag agagg ctaa gactg gtcc tgag	gcaggg ggggca cagttg aattct gcaaga gctgtt cttggg gccagg ttgccc ctcagc	2079 2139 2199 2259 2319 2439 2499 2559 2619 2672

<400> 10

<sup>&</sup>lt;210> 10 <211> 658 <212> PRT <213> Homo Sapien

Met 1	Pro	Val	Ala	Glu 5	Ala	Pro	Gln	Val	Ala 10	Gly	Gly	Gln	Gly	Asp 15	Gly
_	Asp	Gly	Glu 20	Glu	Ala	Glu	Pro	Glu 25		Met	Phe	Lys	Ala 30	Cys	Glu
Asp	Ser	Lуs 35	Arg	Lys	Ala	Arg	Gly 40	Tyr	Leu	Arg	Leu	Val 45	Pro	Leu	Phe
Val	Leu 50	Leu	Ala	Leu	Leu	Val 55	Leu	Ala	Ser	Ala	Gly 60	Val	Leu	Leu	Trp
Tyr 65	Phe	Leu	Gly	Tyr	Lys 70	Ala	Glu	Val	Met	Val 75	Ser	Gln	Val	Tyr	Ser 80
Gly	Ser	Leu	Arg	Val 85	Leu	Asn	Arg	His	Phe 90	Ser	Gln	Asp	Leu	Thr 95	Arg
Arg	Glu	Ser	Ser 100	Ala	Phe	Arg	Ser	Glu 105	Thr	Ala	Lys	Ala	Gln 110	Lys	Met
Leu	Lys	Glu 115	Leu	Ile	Thr	Ser	Thr 120	Arg	Leu	Gly	Thr	Tyr 125	Tyr	Asn	Ser
	130		Tyr			135					140				
145			Val		150					155					160
Lys	Gly	Leu	His	Ser 165	Tyr	Tyr	Asp	Pro	Phe 170	Val	Leu	Ser	Val	Gln 175	Pro
			Gln 180					185					190		
		195	Gly				200					205			
	210		His			215					220				
225			Leu	_	230	_		_		235	_	_			240
			Cys	245					250					255	
_	_		Arg 260					265			_		270		
		275	Gly				280					285			
	290		Val			295					300				
305		_	Glu		310					315					320
			Val	325					330					335	
			Ala 340					345					350		
		355	۷al				360					365			
	370		Cys			375					380	_			_
385		_	Arg		390		-	-		395			_	_	400
			Cys	405					410					415	
		_	Pro 420					425				_	430		_
Gly	Glu	Trp 435	Pro	Trp	Gln	Ala	Ser 440	Leu	Gln	Val	Arg	Gly 445	Arg	His	Ile
_	450	_	Ala			455		_	_		460			_	_
465			Glu	_	470					475		-			480
Leu	Gly	Lys	Val	Trp 485	Gln	Asn	Ser	Arg	Trp 490	Pro	Gly	Glu	Val	Ser 495	Phe

Lys Val S														
70 70	Ser Arg 500	Leu	Leu	Leu	His	Pro 505	Tyr	His	Glu	Glu	Asp 510	Ser	His	
	Asp Val	Ala	Leu	Leu	Gln 520		Asp	His	Pro	Val 525		Arg	Ser	
Ala Ala V 530		Pro	Val	Cys 535		Pro	Ala	Arg	Ser 540		Phe	Phe	Glu	
Pro Gly I	Leu His	Cys	Trp 550		Thr	Gly	Trp	Gly 555		Leu	Arg	Glu	Gly 560	
Gly Pro I	Ile Ser	Asn 565		Leu	Gln	Lys	Val 570		Val	Gln	Leu	Ile 575		
Gln Asp L	Leu Cys 580		Glu	Val	Tyr	Arg 585		Gln	Val	Thr	Pro 590		Met	
Leu Cys A		Tyr	Arg	Lys	Gly 600		Lys	Asp	Ala	Cys 605		Gly	Asp	
Ser Gly G		Leu	Val	Cys 615		Ala	Leu	Ser	Gly 620		Trp	Phe	Leu	
Ala Gly I 625	Leu Val	Ser	Trp 630		Leu	Gly	Cys	Gly 635		Pro	Asn	Tyr	Phe 640	
Gly Val T	Tyr Thr	Arg 645		Thr	Gly	Val	Ile 650		Trp	Ile	Gln	Gln 655		
Val Thr		040					050					000		
<210> 11 <211> 165 <212> DNA <213> Hom <220> <221> CDS <222> (26	A no Sapi													
<223> DNA	A seque:			ling	a ti	cangr	emb	~~~	der					
pro	otease	(MTSI				. 0.1.01		Lane	BCI	rite				
<400> 11 cgcccgggc ttgggactc ctgtcgaat aggggaaca gatgtcaga	ca ggtc cg ggaa ct tcaa aa atgc	agtaa ttato agatt tttct	P-6) ac ac ga ct ac ct ac	protests	ggcct cttgg atgac cctcc cc at	act g tta g gct g gct	atci atci atci agati aagg	ctt gata gcac gagg aa aa	ccgt ctga ctga taga taga at ga	eggte aatge atgete aggte	ege t gga g eg co	ttgt aagta ggcgg ct ga	ogtgga ataaat gagccg	60 120 180 240 294
<400> 11 cgcccgggc ttgggactc ctgtcgaat	ca ggtc cg ggaa ct tcaa aa atgc ag gtcc	agtaa ttatg agatt tttct tgaaa tca	e-6)  ac ac  ga ct  ct ac  g tt  at ac  ttc	prodestgte	ggcct ttgg atgac cctcc Me	act act cas get G. get G. L	tatetategagategaggggggggggggggggggggggg	ctt gata gcac gagg aa aa Lu As	ccgt ctga ctga taga at ga an As	eggte aatge atgete aggte at co sp Pr	ge tae a gga g eg ee ro P:	ttgt aagta ggcgg ct ga co A	egtgga ataaat gagccg et gtt la Val aaa	120 180 240
<pre>&lt;400&gt; 11 cgcccgggc ttgggactc ctgtcgaat aggggaaca gatgtcaga  gaa gcc c Glu Ala E</pre>	ca ggtc. cg ggaa ct tcaa. aa atgc ag gtcc ccc ttc ccc ttc ccc ttc	agtaa ttato agatt tttot tgaaa tca Ser	e-6)  ac ac ga ct ct ac cg tt at ac ttc Phe 15	protest protes	ggcctitggatgaccccat Me tcg Ser	act g tta g caa g gct cg gc et G l ctt Leu gat	ttt pactor	cctt gata gcac gagg aa aa lu As ggc Gly 20 gtt	ccgt ctga ctga taga at ga sn As ctt Leu	eggte aatge aggte at co sp Pr gat Asp	gga	ttgtaagtaggegget geco A.  ttg Leu	egtgga ataaat gagccg et gtt la Val aaa Lys 25 ctg	120 180 240 294
<pre>&lt;400&gt; 11 cgcccgggc ttgggactc ctgtcgaat aggggaaca gatgtcaga  gaa gcc c Glu Ala F 10 ata agt c</pre>	ca ggtca gg ggaa t tcaa aa atgc ag gtcc ccc ttc Pro Phe	agtaa ttato agatt ttot tgaaa tca Ser gca Ala 30	e-6) ac ac ga ct ct ac cg tt at ac ttc Phe 15 cca Pro aag	profestation profe	ggcctcttggatgacctccat Me tcg Ser gca	cact g tta g cac g gct cg gg et G! L ctt Leu gat Asp	tatoral atoral a	gett gata geac gagg aa aa lu As ggc Gly 20 gtt Val	ccgt ctga ctga taga at ga sn As ctt Leu gct Ala	ggtgatggtgat coat Asp	gga	ttgtaagtaggcggct ggc A. ttg Leu atc atc atc	ataaat gagccg ct gtt La Val aaa Lys 25 ctg Leu	120 180 240 294
<pre>&lt;400&gt; 11 cgcccgggc ttgggactc ctgtcgaat aggggaaca gatgtcaga  gaa gcc c Glu Ala F 10 ata agt c Ile Ser F</pre>	ca ggtc. cg ggaa: t tcaa. aa atgc: ag gtcc ccc ttc cro Phe cct gtt cro Val ctg cca Leu Pro 45	agtaa ttato agatt ttot tgaaa tca Ser gca Ala 30 ttg Leu	ttc Phe 15 CCa Pro aag Lys	profest gtt accept acce	tcg Ser gca Ala ttt	act g tta g got g gg t G t ctt Leu gat Asp CCa Pro ggt	ttt Alaa gate agate aagg gg ga y G ttt Phe gct Ala 35 atc	gett gata geac gagg aa aa lu As ggc Gly 20 gtt Val atc Ile	ccgt ctga ctga taga at ga sn As ctt Leu gct Ala gtc Val	ggtgatggtgaggtgaggtgaggtgagggggggggggg	ggt tage of ggt ggt Asp cag Gln ggt ttc	ttgtaagtaggcggct gcco All ttg Leu atc Ile 40 atc Ile gac	ataaat gagccg ct gtt La Val aaa Lys 25 ctg Leu att Ile	120 180 240 294 342

Ala	cga Arg	tgt Cys	gac Asp	gga Gly	gtc Val	tcg Ser	gat Asp	tgc Cys	aaa Lys	Asp	gly aaa	gag Glu	gac Asp	gag Glu	Tyr	582
90	tat	ata	caa	ata	95 ggt	aat	cad	aat	acc	100	ctc	cad	ata	ttc	105	630
Arg	Cys	Val	Arg	Val 110	Gly	Gly	Gln	Asn	Ala 115	Val	Leu	Gln	Val	Phe 120	Thr	030
gct Ala	gct Ala	tcg Ser	tgg Trp 125	aag Lys	acc Thr	atg Met	tgc Cys	tcc Ser 130	gat Asp	gac Asp	tgg Trp	aag Lys	ggt Gly 135	cac His	tac Tyr	678
					gcc Ala											726
					agc Ser											774
					ctc Leu 175											822
cac His	tca Ser	gta Val	tat Tyr	gtg Val 190	agg Arg	gag Glu	gga Gly	tgt Cys	gcc Ala 195	tct Ser	ggc Gly	cac His	gtg Val	gtt Val 200	acc Thr	870
					tgt Cys											918
					tcc Ser											966
					tac Tyr											1014
					gct Ala 255											1062
			Thr	Ile	cag Gln	Val	${ t Gly}$	Leu	Val	Ser	Leu	Leu	Asp		$\mathtt{Pro}$	1110
					gtg Val											1158
					aat Asn											1206
					atg Met											1254
gag	aac	ttc	ccc	gat	gga	aaa	gtg	tgc	tgg	acg	tca	gga	tgg	999	gcc	1302

Glu Asn 330	Phe	Pro	Asp	Gly 335	Lys	Val	Cys	Trp	Thr 340	Ser	Gly	Trp	Gly	Ala 345	
aca gag Thr Glu	gat Asp	gga Gly	ggt Gly 350	gac Asp	gcc Ala	tcc Ser	cct Pro	gtc Val 355	ctg Leu	aac Asn	cac His	gcg Ala	gcc Ala 360	gtc Val	1350
cct ttg Pro Leu															1398
atc atc Ile Ile															1446
gac agc Asp Ser 395	tgc Cys	cag Gln	gjå aaa	gac Asp	agc Ser 400	gl <sup>à</sup> aaa	Gly aaa	ccc Pro	ctg Leu	gtg Val 405	tgt Cys	caa Gln	gag Glu	agg Arg	1494
agg ctg Arg Leu 410															1542
gag gtg Glu Val															1590
tgg atc Trp Ile											tga *	agag	ggaag	333	1639
gataagta	agc (		ga												1656
gataagt: <210> 1: <211> 4: <212> P! <213> He	2 53 RT	cacct				,									1656
<210> 1: <211> 4: <212> P: <213> Ho <400> 1:	2 53 RT Omo S	cacct Sapie	∍n			,		•							1656
<210> 1: <211> 4: <212> P: <213> Ho	2 53 RT Omo S	cacct Sapie	∍n	Pro	Pro	Ala	Val	Glu	Ala	Pro	Phe	Ser	Phe 15	Arg	1656
<210> 1: <211> 4: <212> P: <213> Ho <400> 1: Met Gly	2 53 RT omo s 2 Glu	Sapie Asn Gly	en Asp 5				Lys	10				Ala	15	_	1656
<210> 1: <211> 4: <212> P: <213> Ho <400> 1: Met Gly 1	2 53 RT omo s 2 Glu Phe	Gly	en Asp 5 Leu	Asp	Asp	Leu Ile	Lys 25	10 Ile	Ser	Pro	Val Pro	Ala 30	15 Pro	Asp	1656
<210> 1: <211> 4: <212> P: <213> Ho <400> 1: Met Gly 1 Ser Leu Ala Asp Phe Pro	2 53 RT omo s Glu Phe Ala 35	Sapie Asn Gly 20 Val	Asp 5 Leu Ala	Asp Ala	Asp Gln Gly	Leu Ile 40	Lys 25 Leu	10 Ile Ser	Ser Leu	Pro Leu Ile	Val Pro 45	Ala 30 Leu	15 Pro Lys	Asp Phe	1656
<210> 1: <211> 4: <212> P: <213> Ho <400> 1: Met Gly 1 Ser Leu Ala Asp Phe Pro 50 Ile Gly	2 53 RT omo s Glu Phe Ala 35 Ile	Sapie Asn Gly 20 Val Ile	Asp 5 Leu Ala Val	Asp Ala Ile His	Asp Gln Gly 55	Leu Ile 40 Ile	Lys 25 Leu Ile	10 Ile Ser Ala	Ser Leu Leu Gly	Pro Leu Ile	Val Pro 45 Leu	Ala 30 Leu Ala	15 Pro Lys Leu	Asp Phe Ala Arg	1656
<210> 1: <211> 4: <212> P: <213> Ho <400> 1: Met Gly 1 Ser Leu Ala Asp Phe Pro 50	2 53 RT omo s Glu Phe Ala 35 Ile	Sapie Asn Gly 20 Val Ile Gly	Asp 5 Leu Ala Val Ile Cys	Asp Ala Ile His 70	Asp Gln Gly 55 Phe	Leu Ile 40 Ile Asp	Lys 25 Leu Ile Cys	10 Ile Ser Ala Ser	Ser Leu Leu Gly 75	Pro Leu Ile 60 Lys	Val Pro 45 Leu Tyr	Ala 30 Leu Ala Arg	15 Pro Lys Leu Cys Val	Asp Phe Ala Arg 80	1656
<210> 1: <211> 4: <212> Pi <213> Ho <400> 1: Met Gly 1 Ser Leu Ala Asp Phe Pro 50 Ile Gly 65	2 53 RT omo S 2 Glu Phe Ala 35 Ile Leu	Asn Gly 20 Val Ile Gly Lys Asp	Asp 5 Leu Ala Val Ile Cys 85	Asp Ala Ile His 70 Ile	Asp Gln Gly 55 Phe Glu	Leu Ile 40 Ile Asp Leu	Lys 25 Leu Ile Cys Ile Tyr	10 Ile Ser Ala Ser Ala 90	Ser Leu Leu Gly 75 Arg	Pro Leu Ile 60 Lys Cys	Val Pro 45 Leu Tyr Asp	Ala 30 Leu Ala Arg Gly Val	15 Pro Lys Leu Cys Val 95	Asp Phe Ala Arg 80 Ser	1656
<210> 1: <211> 4: <212> P! <213> Ho <400> 1: Met Gly 1 Ser Leu Ala Asp Phe Pro 50 Ile Gly 65 Ser Ser	2 53 RT omo S 2 Glu Phe Ala 35 Ile Leu Phe Lys	Asn Gly 20 Val Ile Gly Lys Asp	Asp 5 Leu Ala Val Ile Cys 85 Gly	Asp Ala Ile His 70 Ile Glu	Asp Gln Gly 55 Phe Glu Asp	Leu Ile 40 Ile Asp Leu Glu Phe	Lys 25 Leu Ile Cys Ile Tyr 105	10 Ile Ser Ala Ser Ala 90 Arg	Ser Leu Leu Gly 75 Arg Cys	Pro Leu Ile 60 Lys Cys Val	Val Pro 45 Leu Tyr Asp Arg	Ala 30 Leu Ala Arg Gly Val 110	15 Pro Lys Leu Cys Val 95 Gly	Asp Phe Ala Arg 80 Ser Gly	1656
<pre>&lt;210&gt; 1: &lt;211&gt; 4! &lt;212&gt; P! &lt;213&gt; Ho &lt;200&gt; 1: Met Gly 1 Ser Leu Ala Asp Phe Pro 50 Ile Gly 65 Ser Ser Asp Cys Gln Asn Cys Ser</pre>	2 53 RT omo S Glu Phe Ala 35 Ile Leu Phe Lys Ala 115	Asn Gly 20 Val Ile Gly Lys Asp 100 Val	Asp 5 Leu Ala Val Ile Cys 85 Gly Leu	Asp Ala Ile His 70 Ile Glu	Asp Gln Gly 55 Phe Glu Asp Val	Leu Ile 40 Ile Asp Leu Glu Phe 120	Lys 25 Leu Ile Cys Ile Tyr 105 Thr	10 Ile Ser Ala Ser Ala 90 Arg	Ser Leu Leu Gly 75 Arg Cys Ala	Pro Leu Ile 60 Lys Cys Val Ser	Val Pro 45 Leu Tyr Asp Arg Trp 125	Ala 30 Leu Ala Arg Gly Val 110 Lys	15 Pro Lys Leu Cys Val 95 Gly Thr	Asp Phe Ala Arg 80 Ser Gly Met	1656
<pre> &lt;210&gt; 1: &lt;211&gt; 4: &lt;212&gt; P: &lt;213&gt; Ho &lt;213&gt; Ho &lt;400&gt; 1: Met Gly 1 Ser Leu Ala Asp Phe Pro 50 Ile Gly 65 Ser Ser Asp Cys Gln Asn Cys Ser 130 Leu Gly </pre>	2 53 RT DMO S Glu Phe Ala 35 Ile Leu Phe Lys Ala 115 Asp	Asn Gly 20 Val Ile Gly Lys Asp 100 Val Asp	Asp 5 Leu Ala Val Ile Cys 85 Gly Leu Trp	Asp Ala Ile His 70 Ile Glu Gln Lys Tyr	Asp Gln Gly 55 Phe Glu Asp Val Gly 135	Leu Ile 40 Ile Asp Leu Glu Phe 120 His	Lys 25 Leu Ile Cys Ile Tyr 105 Thr	10 Ile Ser Ala Ser Ala 90 Arg Ala	Ser Leu Leu Gly 75 Arg Cys Ala Asn	Pro Leu Ile 60 Lys Cys Val Ser Val 140	Val Pro 45 Leu Tyr Asp Arg Trp 125 Ala	Ala 30 Leu Ala Arg Gly Val 110 Lys	15 Pro Lys Leu Cys Val 95 Gly Thr	Asp Phe Ala Arg 80 Ser Gly Met Gln Ser	1656
<pre>&lt;210&gt; 1: &lt;211&gt; 4: &lt;212&gt; P! &lt;213&gt; Ho &lt;200&gt; 1: Met Gly 1 Ser Leu Ala Asp Phe Pro 50 Ile Gly 65 Ser Ser Asp Cys Gln Asn Cys Ser 130 Leu Gly 145</pre>	2 53 RT DMO S 2 Glu Phe Ala 35 Ile Leu Phe Lys Ala 115 Asp	Asn Gly 20 Val Ile Gly Lys Asp 100 Val Asp Pro	Asp 5 Leu Ala Val Ile Cys 85 Gly Leu Trp Ser	Asp Ala Ile His 70 Ile Glu Gln Lys Tyr 150	Asp Gln Gly 55 Phe Glu Asp Val Gly 135 Val	Leu Ile 40 Ile Asp Leu Glu Phe 120 His Ser	Lys 25 Leu Ile Cys Ile Tyr 105 Thr Tyr	10 Ile Ser Ala Ser Ala 90 Arg Ala Ala	Ser Leu Leu Gly 75 Arg Cys Ala Asn Asn	Pro Leu Ile 60 Lys Cys Val Ser Val 140 Leu	Val Pro 45 Leu Tyr Asp Arg Trp 125 Ala	Ala 30 Leu Ala Arg Gly Val 110 Lys Cys	15 Pro Lys Leu Cys Val 95 Gly Thr Ala	Asp Phe Ala Arg 80 Ser Gly Met Gln Ser 160	1656
<pre> &lt;210&gt; 1: &lt;211&gt; 4: &lt;212&gt; P: &lt;213&gt; Ho &lt;213&gt; Ho &lt;400&gt; 1: Met Gly 1 Ser Leu Ala Asp Phe Pro 50 Ile Gly 65 Ser Ser Asp Cys Gln Asn Cys Ser 130 Leu Gly </pre>	2 53 RT Omo S 2 Glu Phe Ala 35 Ile Leu Phe Lys Ala 115 Asp Phe	Asn Gly 20 Val Ile Gly Lys Asp 100 Val Asp Pro Gln	Asp 5 Leu Ala Val Ile Cys 85 Gly Leu Trp ser Phe 165	Asp Ala Ile His 70 Ile Glu Gln Lys Tyr 150 Arg	Asp Gly 55 Phe Glu Asp Val Gly 135 Val Glu	Leu Ile 40 Ile Asp Leu Glu Phe 120 His Ser Glu	Lys 25 Leu Ile Cys Ile Tyr 105 Thr Tyr Ser	10 Ile Ser Ala Ser Ala Arg Ala Asp Val	Ser Leu Leu Gly 75 Arg Cys Ala Asn 155 Ser	Pro Leu Ile 60 Lys Cys Val Ser Val 140 Leu Ile	Val Pro 45 Leu Tyr Asp Arg Trp 125 Ala Arg	Ala 30 Leu Ala Arg Gly Val 110 Lys Cys Val His	15 Pro Lys Leu Cys Val 95 Gly Thr Ala Ser Leu 175	Asp Phe Ala Arg 80 Ser Gly Met Gln Ser 160 Leu	1656

```
185
Gly Cys Ala Ser Gly His Val Val Thr Leu Gln Cys Thr Ala Cys Gly
       195
               . 200
                                             205
His Arg Arg Gly Tyr Ser Ser Arg Ile Val Gly Gly Asn Met Ser Leu
                      215
                                          220
   210
Leu Ser Gln Trp Pro Trp Gln Ala Ser Leu Gln Phe Gln Gly Tyr His
                230
                                      235
Leu Cys Gly Gly Ser Val Ile Thr Pro Leu Trp Ile Ile Thr Ala Ala
               245
                                  250
                                                      255
His Cys Val Tyr Asp Leu Tyr Leu Pro Lys Ser Trp Thr Ile Gln Val
           260
                            265
Gly Leu Val Ser Leu Leu Asp Asn Pro Ala Pro Ser His Leu Val Glu
       275
                          280
                                           285
Lys Ile Val Tyr His Ser Lys Tyr Lys Pro Lys Arg Leu Gly Asn Asp
                   295
 290
                                      300
Ile Ala Leu Met Lys Leu Ala Gly Pro Leu Thr Phe Asn Glu Met Ile
                  310
                                      315
Gln Pro Val Cys Leu Pro Asn Ser Glu Glu Asn Phe Pro Asp Gly Lys
               325
                                  330
                                                      335
Val Cys Trp Thr Ser Gly Trp Gly Ala Thr Glu Asp Gly Gly Asp Ala
                                                  350
                              345
Ser Pro Val Leu Asn His Ala Ala Val Pro Leu Ile Ser Asn Lys Ile
       355
                           360
                                              365
Cys Asn His Arg Asp Val Tyr Gly Gly Ile Ile Ser Pro Ser Met Leu
                     375
                                        380
 370
Cys Ala Gly Tyr Leu Thr Gly Gly Val Asp Ser Cys Gln Gly Asp Ser
                  390
                                      395
Gly Gly Pro Leu Val Cys Gln Glu Arg Arg Leu Trp Lys Leu Val Gly
               405
                                  410
                                                  415
Ala Thr Ser Phe Gly Ile Gly Cys Ala Glu Val Asn Lys Pro Gly Val
           420
                              425
                                                 430
Tyr Thr Arg Val Thr Ser Phe Leu Asp Trp Ile His Glu Gln Met Glu
                    440
       435
                                             445
Arg Asp Leu Lys Thr
  450
<210> 13
<211> 23
<212> DNA
<213> Artificial Sequence
<223> Oligonucleotide Primer
<221> misc_feature
<222> (0) ... (0)
<223> N= Inosine
<400> 13
tggrtnvtnw sngcnrcnca ytg
<210> 14
<211> 30
<212> DNA
<213> Artificial Sequence
<220>
```

<223> Oligonucleotide Primer

<221> misc\_feature <222> (0)...(0) <223> N= Inosine

<400> nggnco	14 encen swrtenecyt nreanghrte	30
<210><211><212><213>	30	
<220> <223>	Oligonucleotide primer	
<400> tcaćco	15 gagaa gatgatgtgt gcaggcatcc	30
<210><211><211><212><213>	29	
<220> <223>	Oligonucleotide primer	
<400> gggaca	16 agggg ctgtaaggca gggaatgag	29
<210><211><211><212><213>	25	
<220> <223>	Oligonucleotide primer	
<400> cccgca	17 agcca tagccccagc taacg	25
<210><211><211><212><213>	27	
<400> gcagac	18 gatg cgtaccaggg ggaagtc	27
<210><211><211><212><213>	39	
<220> <223>	Oligonucleotide primer	
<400> ctcgag	19 gaaaa gagtggtggg tggggaggag gcctctgtg	39
<210><211><212><213>	30	
<220> <223>	Oligonucleotide primer	

<400> gcggcd		30
<210> <211> <212> <213>	27	
<220> <223>	Oligonucleotide Primer	
<400> cctcca	21 acggt gctgtggacc gtgttcc	27
<210> <211> <212> <213>	24	
<220> <223>	Oligonucleotide Primer	
<400> cctcgd	22 egcaa ggegeeecag eeeg	24
<210> <211> <212> <213>	32	
<220> <223>	Oligonucleotide Primer	
<400> gcgtgg		32
<210><211><212><213>	27	
<220> <223>	Oligonucleotide Primer	
<400>	24 acggt gctgtggacc gtgttcc	27
<210><211><211><212><213>	24	
<220> <223>	Oligonucleotide Primer	
<400>	25 egcaa ggegeeceag eeeg	24
<210><211><212><213>	26	

<220> <223> Oligonucleotide Primer	
<400> 26 tcatcggcca gagggtgatc agtgag	26
<210> 27 <211> 28 <212> DNA <213> Artificial Sequence	
<220> <223> Oligonucleotide Primer	
<400> 27 cctcctcagt gcataggcat caaaccag	28
<210> 28 <211> 42 <212> DNA <213> Artificial Sequence	
<220> <223> Oligonucleotide Primer	
<400> 28 tctctcgaga aaagaattgt tggtggagct gtgtcctccg ag	42
<210> 29 <211> 31 <212> DNA <213> Artificial Sequence	
<220> <223> Oligonucleotide Primer	
<400> 29 aggtgggcct tgctttgcag gggggcagtt c	31
<210> 30 <211> 26 <212> DNA <213> Artificial Sequence	
<220> <223> Oligonucleotide Primer	
<400> 30 tcacgcatcg tgggtggaac atgtcc	26
<210> 31 <211> 26 <212> DNA <213> Artificial Sequence	
<220> <223> Oligonucleotide Primer	
<400> 31 acccacctcc atctgctcgt ggatcc	26
<210> 32	

<211> <212> <213>		
<220> <223>	Oligonucleotide primer	
<400> ccacaç	32 geete etetettgae acaccag	27
<210><211><212><213>	26	
<220> <223>	Oligonucleotide primer	
<400> acgcco	33 ectgt ggatcatcac tgctgc	26
<210><211><211><212><213>	27	
<220> <223>	Oligonucleotide primer	
<400> tccct	34 ccctc acatatactg agtggtg	27
<210><211><211><212><213>	26	
<220> <223>	Oligonucleotide primer	
<400> cgacto	35 getea gggaagteag atgteg	26
<210><211><212><213>	36	
<220> <223>	Oligonucleotide primer	
<400> gcggcd	36 egcac tataceccag tgttetettt gateca	36
<210><211><211><212><213>	27	
<220> <223>	Oligonucleotide primer	

<400> 37 ctggtgtgtc aagagaggag gctgtgg	27
<210> 38 <211> 28 <212> DNA <213> Artificial Sequence	
<220> <223> Oligonucleotide primer	
<400> 38 actcaggtgg ctacttatec cetteete	28
<210> 39 <211> 42 <212> DNA <213> Artificial Sequence	
<220> <223> Oligonucleotide Primer	
<400> 39 tctctcgaga aaagagtggt gggtggggag gaggcctctg tg	42
<210> 40 <211> 34 <212> DNA <213> Artificial Sequence	
<220> <223> Oligonucleotide Primer	
<400> 40 attcgcggcc gcattacagc tcagccttcc agac	34
<210> 41 <211> 42 <212> DNA <213> Artificial Sequence	
<220> <223> Oligonucleotide Primer	
<400> 41 tctctcgaga aaagaattgt tggtggagct gtgtcctccg ag	42
<210> 42 <211> 37 <212> DNA <213> Artificial Sequence	
<220> <223> Oligonucleotide Primer	
<400> 42 attcgcggcc gctcaggtca ccacttgctg gatccag	37
<210> 43 <211> 36 <212> DNA <213> Artificial Sequence	

<220> <223> Oligonucleotide	Primer				
<400> 43 ctcgagaaac gcatcgtggg	tggaaacatg	tccttg			36
<210> 44 <211> 28 <212> DNA <213> Artificial Seque	ence				
<220> <223> Oligonucleotide	Primer				
<400> 44 actcaggtgg ctacttatcc	ccttcctc				28
<210> 45 <211> 9276 <212> DNA <213> Pichia pastoris					
<400> 45					
agatctaaca tccaaagacg gtccattctc acacataagt	gccaaacgca	acaggagggg	atacactagc	agcagaccgt	60 120
tgcaaacgca ggacctccac agcccagtta ttgggcttga					180 240
acaccatgac tttattagcc	tgtctatcct	ggcccccctg	gcgaggttca	tgtttgttta	300
tttccgaatg caacaagctc	cgcattacac	ccgaacatca	ctccagatga	gggctttctg	360
agtgtggggt caaatagttt					420
gtcttggaac ctaatatgac					480 540
ttgaaatgct aacggccagt cttgtttggt attgattgac					600
ctctatcgct tctgaacccc					660
ttttggatga ttatgcattg					720
gctgatagcc taacgttcat					780
atataaacag aaggaagctg					840
actttcataa ttgcgactgg caacttgaga agatcaaaaa					900 960
tcaattttta ctgcagtttt					1020
acaacagaag atgaaacggc					1080
gaaggggatt tcgatgttgc	tgttttgcca	ttttccaaca	gcacaaataa	cgggttattg	1140
tttataaata ctactattgc	cagcattgct	gctaaagaag	aaggggtatc	tctcgagaaa	1200
agagaggetg aagettaegt					1260 1320
atgactgttc ctcagttcaa tcaagaggat gtcagaatgc					1380
ttatttqtaa cctatataqt					1440
ttgctcctga tcagcctatc					1500
tcattcgagt ttgatgtttt					1560
agtgagaagt tcgtttgtgc					1620
taaattgcta acgcagtcag tcggcaccgt caccctggat					1680 1740
tettqeqqqa tateqtecat					1800
tatatgcgtt gatgcaattt					1860
gccgccgccc agtcctgctc	gcttcgctac	ttggagccac	tatcgactac	gcgatcatgg	1920
cgaccacacc cgtcctgtgg					1980
taaataagtc ccagtttctc					2040 2100
tcaacagcag ccagatccat tcttgtgaag tgatgaactt					2160
tatccgtacg ttggcaaagt					2220
gagtaggcac caacaacac	agatccagcg	tgttgtactt	gatcaacata	agaagaagca	2280
ttctcgattt gcaggatcaa	gtgttcagga	gcgtactgat	tggacatttc	caaagcctgc	2340

2400 tegtaggttg caacegatag ggttgtagag tgtgcaatac acttgegtac aatttcaace cttggcaact gcacagettg gttgtgaaca gcatetteaa ttetggcaag eteettgtet 2460 gtcatatcga cagccaacag aatcacctgg gaatcaatac catgttcagc ttgagacaga aggtctgagg caacgaaatc tggatcagcg tatttatcag caataactag aacttcagaa 2520 2580 2640 ggcccagcag gcatgtcaat actacacagg gctgatgtgt cattttgaac catcatcttg gcagcagtaa cgaactggtt tcctggacca aatattttgt cacacttagg aacagtttct 2700 gttccgtaag ccatagcagc tactgcctgg gcgcctcctg ctagcacgat acacttagca ccaaccttgt gggcaacgta gatgacttct ggggtaaggg taccatcctt cttaggtgga gatgcaaaaa caatttcttt gcaaccagca actttggcag gaacacccag catcagggaa gtggaaggca gaattgcggt tccaccagga atatagaggc caactttctc aataggtctt 2760 2820 2880 2940 3000 gcaaaacgag agcagactac accagggcaa gtctcaactt gcaacgtctc cgttagttga getteatgga attteetgae gttatetata gagagateaa tggetetett aaegttatet 3060 ggcaattgca taagttctc tgggaaagga gcttctaaca caggtgtctt caaagcgact ccatcaaact tggcagttag ttctaaaagg gctttgtcac cattttgacg aacattgtcg acaattggtt tgactaattc cataatctgt tccgttttct ggataggacg acgaagggca 3120 3180 3240 3300 tottcaattt ottgtgagga ggoottagaa acgtcaattt tgcacaattc aatacgacct tcagaaggga cttctttagg tttggattct tctttaggtt gttccttggt gtatcctggc ttggcatctc ctttccttct agtgaccttt agggacttca tatccaggtt tctctccacc 3360 3420 tegtecaacg teacacegta citggeacat ciaactaatg caaaataaaa taagteagea 3480 catteceagg ctatatette ettggattta gettetgeaa gttcateage tteeteeeta 3540 attttagcgt tcaacaaaac ttcgtcgtca aataaccgtt tggtataaga accttctgga 3600 gcattgctct tacgatccca caaggtggct tccatggctc taagaccctt tgattggcca 3660 aaacaggaag tgcgttccaa gtgacagaaa ccaacacctg tttgttcaac cacaaatttc 3720 aagcagtete cateacaate caattegata cecageaact tttgagttge tecagatgta 3780 gcaccettat accacaacc gtgacgacga gattggtaga ctccagtttg tgtccttata 3840 gcctccggaa tagacttttt ggacgagtac accaggccca acgagtaatt agaagagtca 3900 gccaccaaag tagtgaatag accatcgggg cggtcagtag tcaaagacgc caacaaaatt tcactgacag ggaacttttt gacatcttca gaaagttcgt attcagtagt caattgccga 3960 4020 gcatcaataa tggggattat accagaagca acagtggaag tcacatctac caactttgcg 4080 4140 gtctcagaaa aagcataaac agttctacta ccgccattag tgaaactttt caaatcgccc agtggagaag aaaaaggcac agcgatacta gcattagcgg gcaaggatgc aactttatca accagggtcc tatagataac cctagcgcct gggatcatcc tttggacaac tctttctgcc aaatctaggt ccaaaatcac ttcattgata ccattattgt acaacttgag caagttgtcg 4200 4260 4320 atcagetect caaattggte etetgtaaeg gatgaeteaa ettgeaeatt aacttgaage 4380 4440 tcagtcgatt gagtgaactt gatcaggttg tgcagctggt cagcagcata gggaaacacg getttteeta ecaaacteaa ggaattatea aaetetgeaa eaettgegta tgeaggtage 4500 aagggaaatg tcatacttga agtcggacag tgagtgtagt cttgagaaat tctgaagccg 4560 tatttttatt atcagtgagt cagtcatcag gagatcctct acgccggacg catcgtggcc 4620 gacctgcagg ggggggggg gcgctgaggt ctgcctcgtg aagaaggtgt tgctgactca 4680 4740 taccaggeet gaategeece ateatecage cagaaagtga gggageeaeg gttgatgaga gctttgttgt aggtggacca gttggtgatt ttgaactttt gctttgccac ggaacggtctgcgg gaagatgcgt gatctgatcc ttcaactcag caaaagttcg atttattcaa 4800 4860 caaagccgcc gtcccgtcaa gtcagcgtaa tgctctgcca gtgttacaac caattaacca 4920 attctgatta gaaaaactca tcgagcatca aatgaaactg caatttattc atatcaggat 4980 5040 tatcaatacc atatttttga aaaagccgtt tctgtaatga aggagaaaac tcaccgaggc agttccatag gatggcaaga tcctggtatc ggtctgcgat tccgactcgt ccaacatcaa tacaacctat taatttcccc tcgtcaaaaa taaggttatc aagtgagaaa tcaccatgag 5100 5160 tgacgactga atccggtgag aatggcaaaa gcttatgcat ttctttccag acttgttcaa 5220 5280 caggccagcc attacgctcg tcatcaaaat cactcgcatc aaccaaaccg ttattcattc gtgattgege etgagegaga egaaataege gategetgtt aaaaggacaa ttacaaacag 5340 gaatcgaatg caaccggcgc aggaacactg ccagcgcatc aacaatattt tcacctgaat caggatattc ttctaatacc tggaatgctg ttttcccggg gatcgcagtg gtgagtaacc 5400 5460 atgcatcatc aggagtacgg ataaaatgct tgatggtcgg aagaggcata aattccgtca 5520 5580 gccagtttag tctgaccatc tcatctgtaa catcattggc aacgctacct ttgccatgtt 5640 tcagaaacaa ctctggcgca tcgggcttcc catacaatcg atagattgtc gcacctgatt gcccgacatt atcgcgagcc catttatacc catataaatc agcatccatg ttggaattta 5700 atcgcggcct cgagcaagac gtttcccgtt gaatatggct cataacaccc cttgtattac 5760 tgtttatgta agcagacagt tttattgttc atgatgatat atttttatct tgtgcaatgt 5820 5880 aacatcagag attttgagac acaacgtggc tttccccccc ccccetgcag gtcggcatca 5940 ccggcgccac aggtgcggtt gctggcgcct atatcgccga catcaccgat ggggaagatc gggctcgcca cttcgggctc atgagcgctt gtttcggcgt gggtatggtg gcaggccccg tggccggggg actgttgggc gccatctcct tgcatgcacc attccttgcg gcggcggtgc 6000 6060

	tcaacggcct	caacctacta	ctgggctgct	tcctaatgca	ggagtcgcat	aagggagagc		6120
	gtcgagtatc	tatgattgga	agtatgggaa	tggtgatacc	cgcattcttc	agtgtcttga		6180
			ccaactaaag					6240
			gtagactcga					6300
			gtaaatgaag					6360
	gaacaaactt	cttgtttcga	actttttcgg	tgccttgaac	tataaaatgt	agagtggata		6420
			ggcaaatgct					6480
	gtttgtagat	actgatgcca	acttcagtga	caacgttgct	atttcgttca	aaccattccg		6540
			gtttgtctac					6600
			tttgaggtca					6660
			aggcgataaa				*	6720
			ctgtattaaa					6780
			tgttttagag					6840
			gaaatttatc					6900
			acatgcagct					6960
			cccgtcaggg					7020
			gtagcgatag					7080
,	tcagagcaga	ttgtactgag	agtgcaccat	atgcggtgtg	aaataccgca	cagatgcgta	•	7140
	aggagaaaat	accgcatcag	gcgctcttcc	gcttcctcgc	tcactgactc	getgegeteg		7200
			ggtatcagct					7260
			aaagaacatg					7320
			ggcgtttttc					7380
	aaaaatcgac	gctcaagtca	gaggtggcga	aacccgacag	gactataaag	ataccaggcg		7440
			cgtgcgctct					7500
			gggaagcgtg					7560
			tcgctccaag					7620
			cggtaactat					7680
			cactggtaac					7740
						agtatttggt		7800 7860
			agttaccttc					7920
			cggtggtttt					7980
			tcctttgatc					8040
	gaaaacccac	gccaagggac	tttggtcatg	agattattaa	tatataaata	acctagate		8100
	gagagttagg	aaaaatyaay	ttttaaatca cagtgaggca	acctatataa	catatatat	atttcattca		8160
								8220
	tccatagttg ggccccagtg							8280
J.	atasaggag	gaggggaaga	ggccgagcgc	agaagtggtc	ctccaga	atccccctcc		8340
	ataaaccagc	ttaattatta	ggccgagcgc	agaageggee	attaccaat	taatagtttg		8400
• '			tgcaggcatc					8460
	tcattcacct	ccgcttccca	acgatcaagg	caaattacat	gatccccat	attatacasa		8520
•	aaaacaatta	actacttaa	tcctccgatc	attatcagaa	gtaagttggc	cacaatatta		8580
- '			actgcataat					8640
			ctcaaccaag					8700
			aacacgggat					8760
	atactcatca	ttagaaaaca	ttcttcgggg	cgaaaact.ct	caaggatett	accactatta		8820
			cactcgtgca					8880
	accagcottt	ctagataaac	aaaaacagga	aggcaaaatg	ccccaaaaaa	gggaataagg		8940
			actcatactc					9000
			cggatacata					9060
			ccgaaaagtg					9120
	atgacattaa	cctataaaaa	taggcgtatc	acgaggccct	ttcatcttca	agaattaatt		9180
			atcgataagc					9240
			taacagttat		555-5	5 5 -		9276
	J 1,555	<u> </u>	<b>9</b>	<b>J</b>				

agegeecaat aegeaaaceg ceteteeceg egegttggee gatteattaa tgeagetgge

<sup>&</sup>lt;210> 46 <211> 3908 <212> DNA

<sup>&</sup>lt;213> Escherichia coli

120 acgacaggtt tcccgactgg aaagcgggca gtgagcgcaa cgcaattaat gtgagttagc tcactcatta ggcaccccag gctttacact ttatgcttcc ggctcgtatg ttgtgtggaa 180 240 ttgtgagcgg ataacaattt cacacaggaa acagctatga ccatgattac gccaagcttg gtaccgaget eggatecaet agtaaeggee geeagtgtge tggaattege cettaaggge 300 gaattotgoa gatatocato acactggogg cogetogago atgoatotag agggoocaat 360 420 tegecetata gtgagtegta ttacaattea etggeegteg ttttacaaeg tegtgaetgg 480 gaaaaccctg gcgttaccca acttaatcgc cttgcagcac atcccccttt cgccagctgg cgtaatagcg aagaggcccg caccgatcgc ccttcccaac agttgcgcag cctgaatggc gaatgggacg cgcctgtag cggcgcatta agcgcggcgg gtgtggtggt tacgcgcagc gtgaccgcta cacttgccag cgccctagcg cccgctcctt tcgctttctt cccttctt 540 600 660 ctcgccacgt tcgccggctt tccccgtcaa gctctaaatc gggggctccc tttagggttc 720 780 cgatttagag ctttacggca cctcgaccgc aaaaaacttg atttgggtga tggttcacgt agtgggccat cgccctgata gacggttttt cgccctttga cgttggagtc cacgttcttt 840 900 aatagtggac tettgtteca aactggaaca acactcaacc etategeggt etattetttt gatttataag ggattttgcc gatttcggcc tattggttaa aaaatgagct gatttaacaa 960 attcagggcg caagggctgc taaaggaacc ggaacacgta gaaagccagt ccgcagaaac 1020 1080 ggtgctgacc ccggatgaat gtcagctact gggctatctg gacaagggaa aacgcaagcg caaagagaaa gcaggtagct tgcagtgggc ttacatggcg atagctagac tgggcggttt 1140 1200 tatggacagc aagcgaaccg gaattgccag ctggggcgcc ctctggtaag gttgggaagc cctgcaaagt aaactggatg gctttcttgc cgccaaggat ctgatggcgc aggggatcaa 1260 1320 gatctgatca agagacagga tgaggatcgt ttcgcatgat tgaacaagat ggattgcacg caggttetee ggeegettgg gtggagagge tatteggeta tgactgggea caacagacaa 1380 teggetgete tgatgeegee gtgtteegge tgteagegea ggggegeeeg gttetttttg 1440 1500 tcaagaccga cctgtccggt gccctgaatg aactgcagga cgaggcagcg cggctatcgt ggctggccac gacgggcgtt ccttgcgcag ctgtgctcga cgttgtcact gaagcgggaa 1560 gggactggct gctattgggc gaagtgccgg ggcaggatct cctgtcatct cgccttgctc 1620 ctgccgagaa agtatccatc atggctgatg caatgcggcg gctgcatacg cttgatccgg ctacctgcc attcgaccac caagcgaaac atcgcatcga gcgagcacgt actcggatgg 1680 1740 aagccggtct tgtcgatcag gatgatctgg acgaagagca tcagggggctc gcgccagccg 1800 aactgttege caggetcaag gegegeatge eegaeggega ggatetegte gtgatecatg 1860 1920 gcgatgcctg cttgccgaat atcatggtgg aaaatggccg cttttctgga ttcaacgact gtggccggct gggtgtggcg gaccgctatc aggacatagc gttggatacc cgtgatattg ctgaagagct tggcggcgaa tgggctgacc gcttcctcgt gctttacggt atcgccgctc ccgattcgca gcgcatcgcc ttctatcgcc ttcttgacga gttcttctga attgaaaaag 1980 2040 2100 2160 gaagagtatg agtattcaac atttccgtgt cgcccttatt cccttttttg cggcattttg ccttcctgtt tttgctcacc cagaaacgct ggtgaaagta aaagatgctg aagatcagtt 2220 gggtgcacga gtgggttaca tcgaactgga tctcaacagc ggtaagatcc ttgagagttt tcgccccgaa gaacgttttc caatgatgag cacttttaaa gttctgctat gtcatacact 2280 2340 attatcccgt attgacgccg ggcaagagca actcggtcgc cgggcgcggt attctcagaa 2400 tgacttggtt gagtactcac cagtcacaga aaagcatctt acggatggca tgacagtaag 2460 agaattatgc agtgctgcca taaccatgag tgataacact gcggccaact tacttctgac aacgatcgga ggaccgaagg agctaaccgc ttttttgcac aacatggggg atcatgtaac 2520 2580 togoottgat ogttgggaac oggagotgaa tgaagocata ocaaacgacg agagtgacac-2640 cacgatgcct gtagcaatgc caacaacgtt gcgcaaacta ttaactggcg aactacttac 2700 tctagcttcc cggcaacaat taatagactg gatggaggcg gataaagttg caggaccact tctgcgctcg gcccttccgg ctggctggtt tattgctgat aaatctggag ccggtgagcg 2760 2820 tgggtctcgc ggtatcattg cagcactggg gccagatggt aagccctccc gtatcgtagt tatctacacg acggggagtc aggcaactat ggatgaacga aatagacaga tcgctgagat 2880 2940 aggtgcctca ctgattaagc attggtaact gtcagaccaa gtttactcat atatacttta 3000 gattgattta aaacttcatt tttaatttaa aaggatctag gtgaagatcc tttttgataa tctcatgacc aaaatccctt aacgtgagtt ttcgttccac tgagcgtcag accccgtaga 3060 3120 aaagatcaaa ggatcttctt gagatccttt ttttctgcgc gtaatctgct gcttgcaaac aaaaaaacca ccgctaccag cggtggtttg tttgccggat caagagctac caactctttt 3180 3240 tccgaaggta actggcttca gcagagcgca gataccaaat actgtccttc tagtgtagcc 3300 3360 gtagttaggc caccacttca agaactctgt agcaccgcct acatacctcg ctctgctaat cctgttacca gtggctgctg ccagtggcga taagtcgtgt cttaccgggt tggactcaag 3420 acgatagtta ccggataagg cgcagcggtc gggctgaacg gggggttcgt gcacacagcc cagcttggag cgaacgacct acaccgaact gagataccta cagcgtgagc attgagaaag 3480 3540 cgccacgctt cccgaaggga gaaaggcgga caggtatccg gtaagcggca gggtcggaac 3600 3660 aggagagege aegagggage tteeaggggg aaaegeetgg tatetttata gteetgtegg gtttcgccac ctctgacttg agcgtcgatt tttgtgatgc tcgtcagggg ggcggagcct 3720 atggaaaaac gccagcaacg cggccttttt acggttcctg gccttttgct ggccttttgc 3780

```
teacatqtte ttteetqcqt tateceetqa ttetqtqqat aaccqtatta cegcetttga
                                                                                      3840
                                                                                      3900
  qtqaqctqat accqctcqcc qcaqccgaac gaccgagcgc agcgagtcag tgagcgagga
                                                                                      3908
  agcggaag
  <210> 47
  <211> 46
  <212> DNA
  <213> Artificial Sequence
  <220>
 <223> Oligonucleotide primer
  <400> 47
                                                                                         46
  ggaattccat atgccgcgct ttaaagtggt gggtggggag gaggcc
  <211> 32
  <212> DNA
  <213> Artificial Sequence
  <223> Oligonucleotide primer
  <400> 48
                                                                                         32
  cgcgataccc gttacagctc agccttccag ac
  <210> 49
  <211> 3147
  <212> DNA
  <213> Homo Sapien
  <220>
  <221> CDS
  <222> (1865)...(2590)
  <223> Nucleic acid sequence of protease domain of MTSP1
  <400> 49
  tcaagagegg cctcggggta ccatggggag cgatcgggcc cgcaagggcg gaggggccc
                                                                                       60
  gaaggacttc ggcgcgggac tcaagtacaa ctcccggcac gagaaagtga atggcttgga
                                                                                      120
                                                                                      180
 ggaaggegtg gagtteetge cagteaacaa egteaagaag gtggaaaage atggeeeggg
  gegetgggtg gtgetggeag cegtgetgat eggeeteete ttggtettge tggggategg etteetggtg tggeatttge agtaceggga egtgegtgte cagaaggtet teaatggeta
                                                                                      240
                                                                                      300
catgaggatc acaaatgaga attttgtgga tgcctacgag aactccaact ccactgagtt
                                                                                      360
  tqtaaqcctq qccaqcaaqq tqaaggacgc gctgaagctg ctgtacagcg gagtcccatt
                                                                                      420
                                                                                      480
 cctgggcccc taccacaagg agtcggctgt gacggccttc agcgagggca gcgtcatcgc
 ctactactgg tetgagttea geatecegea geacetggtg gaggaggeeg agegegteat ggeegaggag egegtagtea tgetgeecee gegggegege teeetgaagt cetttgtggt eaceteagtg gtggetttee ceaeggaete caaaacagta cagaggaece aggacaacag
                                                                                      540
                                                                                      600
                                                                                      660
  ctgcagcttt ggcctgcacg cccgcggtgt ggagctgatg cgcttcacca cgcccggctt
                                                                                      720
                                                                                      780
ceetgacage ceetaceeeg eteatgeeeg etgecagtgg geeetgeggg gggacgeega
ctcagtgetg agectcacet teegeagett tgacettgeg teetgegaeg agegeggeag egacetggtg aeggtgtaca acaceetgag ecceatggag ecceaegece tggtgeagtt gtgtggeaec taccetecet ectacaacet gacetteeae teeteceaga aegteetget
                                                                                      840
                                                                                      900
                                                                                      960
  catcacactg ataaccaaca ctgagcggcg gcatcccggc tttgaggcca ccttcttcca
                                                                                     1020
                                                                                     1080
  gctgcctagg atgagcagct gtggaggccg cttacgtaaa gcccagggga cattcaacag
  cocctactac ccaggocact acccacccaa cattgactgo acatggaaca ttgaggtgco
                                                                                     1140
  caacaaccag catgtgaagg tgagcttcaa attcttctac ctgctggagc ccggcgtgcc tgcgggcacc tgccccaagg actacgtgga gatcaatggg gagaaatact gcggagagag
                                                                                     1200
                                                                                     1260
  gtcccagttc gtcgtcacca gcaacagcaa caagatcaca gttcgcttcc actcagatca
                                                                                     1320
                                                                                     1380
  gtcctacacc gacaccggct tcttagctga atacctctcc tacgactcca gtgacccatg
  cccggggcag ttcacgtgcc gcacggggcg gtgtatccgg aaggagctgc gctgtgatgg
                                                                                     1440
                                                                                     1500
  ctgggccgac tgcaccgacc acagcgatga gctcaactgc agttgcgacg ccggccacca
```

## WO 01/57194 PCT/US01/03471 36/66

gttcacgtgc aagaacaagt tctgcaagcc cctcttctgg gtctgcgaca gtgtgaacga ctgcggagac aacagcgacg agcagggtg cagttgtccg gcccagacct tcaggtgttc caatgggaag tgcctctcga aaagccagca gtgcaatggg aaggacgact gtggggacgg gtccgacgag gctcctgcc ccaaggtgaa cgtcgtcact tgtaccaaac acacctaccg ctgcctcaat gggctctgct tgagcaaggg caaccctgag tgtgacgga aggaggactg tagcgacggc tcagatgaga aggactgcga ctgtgggctg cggtcattca cgagacaggc tcgt gtt gtt ggg ggc acg gat gcg gat gag ggc gag tgg ccc tgg cag  Val Val Gly Gly Thr Asp Ala Asp Glu Gly Glu Trp Pro Trp Gln  1 5 10 15													1560 1620 1680 1740 1800 1860 1909			
gta Val	agc Ser	ctg Leu	cat His	gct Ala 20	ctg Leu	ggc Gly	cag Gln	ggc Gly	cac His 25	atc Ile	tgc Cys	ggt Gly	gct Ala	tcc Ser 30	ctc Leu	1957
						gtc Val										2005
						gac Asp										2053
						cgc Arg 70										2101
						cac His										2149
						gag Glu										2197
						ctg Leu										2245
ggc	aag Lys	gcc Ala 130	atc Ile	tgg Trp	gtc Val	acg Thr	ggc Gly 135	tgg Trp	gga Gly	cac His	acc Thr	cag Gln 140	tat Tyr	gga Gly	ggc Gly	2293
act Thr	ggc Gly 145	gcg Ala	ctg Leu	atc Ile	ctg Leu	caa Gln 150	aag Lys	ggt Gly	gag Glu	atc Ile	cgc Arg 155	gtc Val	atc Ile	aac Asn	cag Gln	2341
						ctg Leu										2389
tgc Cys	gtg Val	ggc Gly	ttc Phe	ctc Leu 180	agc Ser	ggc Gly	ggc Gly	gtg Val	gac Asp 185	tcc Ser	tgc Cys	cag Gln	ggt Gly	gat Asp 190	tcc Ser	2437
gly aaa	gga Gly	ccc Pro	ctg Leu 195	tcc Ser	agc Ser	gtg Val	gag Glu	gcg Ala 200	gat Asp	gly aaa	cgg Arg	atc Ile	ttc Phe 205	cag Gln	gcc Ala	2485
						gac Asp										2533

PCT/US01/03471

gtg tac aca agg Val Tyr Thr Arg 225										
ggg gta tag gggd Gly Val * 240	ccggggc cacco	caaatg tgtacad	cctg cggggccacc	2630						
catcgtccac cccagtgtgc acgcctgcag gctggagact ggaccgctga ctgcaccagc gcccccagaa catacactgt gaactcaatc tccagggctc caaatctgcc tagaaaacct ctcgcttcct cagcctccaa agtggagctg ggaggtagaa ggggaggaca ctggtggttc tactgaccca actggggca aaggtttgaa gacacagcct cccccgccag ccccaagctg ggccgaggcg cgtttgtgta tatctgcac ccctgtctgt aaggagcag gggaacggag cttcggagcc tcctcagtga aggtggtggg gctgccggat ctgggctgtg gggcccttgg gcaccctct tgaggaagcc caggctcgga ggaccctgga aaacagacg gtctgagact gaaattgtt taccagctcc cagggtggac ttcagtgtgt gtatttgtgt aaatgggtaa aacaatttat ttcttttaa aaaaaaaaa aaaaaaa										
<210> 50 <211> 241 <212> PRT <213> Homo Sapien										
<400> 50 Val Val Gly Gly 1	Thr Asp Ala	Asp Glu Gly (	Glu Trp Pro Trp	Gln Val 15						
Ser Leu His Ala	Leu Gly Gln	Gly His Ile (	Cys Gly Ala Ser	Leu Ile						
Ser Pro Asn Trp	Leu Val Ser	Ala Ala His (	Cys Tyr Ile Asp 45	Asp Arg						
Gly Phe Arg Tyr 50	Ser Asp Pro 55	Thr Gln Trp	Thr Ala Phe Leu 60	Gly Leu						
His Asp Gln Ser 65	Gln Arg Ser 70		Val Gln Glu Arg 75	Arg Leu 80						
Lys Arg Ile Ile	Ser His Pro 85	Phe Phe Asn A	Asp Phe Thr Phe	Asp Tyr 95						
Asp Ile Ala Leu 100	Leu Glu Leu	Glu Lys Pro A 105	Ala Glu Tyr Ser 110	Ser Met						
Val Arg Pro Ile 115	Cys Leu Pro	Asp Ala Ser I 120	His Val Phe Pro 125	Ala Gly						
Lys Ala Ile Trp 130	Val Thr Gly 135	Trp Gly His	Thr Gln Tyr Gly 140	Gly Thr						
Gly Ala Leu Ile 145	Leu Gln Lys 150		Arg Val Ile Asn 155	Gln Thr 160						
Thr Cys Glu Asn	Leu Leu Pro 165	Gln Gln Ile 1	Thr Pro Arg Met	Met Cys 175						
Val Gly Phe Leu 180	Ser Gly Gly	Val Asp Ser (	Cys Gln Gly Asp 190	Ser Gly						
Gly Pro Leu Ser 195	Ser Val Glu		Arg Ile Phe Gln 205	Ala Gly						
Val Val Ser Trp 210	Gly Asp Gly 215			Gly Val						
Tyr Thr Arg Leu 225 Val				Thr Gly 240						

<sup>&</sup>lt;210> 51 <211> 46 <212> DNA <213> Artificial Sequence

<220> <223>	Oligonucleoide Primer	
<400> tctctc	51 gaga aaagagtggt gggtgggtgg ggaggaggcc tctgtg	46
<210><211><212><213>	43	
<400> gctcct	52 catc aaagaagggc agagagatgg gcctgactgt gcc	43
<210><211><211><212><213>	34	
<220> <223>	Oligonucleoide Primer	
<400> attcgc	53 eggec geattacage teageettee agae	34
<210><211><212><213>	43	
<220> <223>	Oligonucleoide Primer	
<400> ggcaca	54 agtca ggcccatctc tctgcccttc tttgatgagg agc	43
<210><211><211><212><213>	28	
<220> <223>	Oligonucleoide Primer	
<400>	55 ettet teaatgaett caeetteg	28
<210> <211> <212> <213>	18	
<220> <223>	Oligonucleoide Primer	
<400> taccto	56 etect aegaetee	18
<210><211><212><212><213>	25	

<220> <223> Oligonucleoide Primer <400> 57 gaggttctcg caggtggtct ggttg 25 <210> 58 <211> 39 <212> DNA <213> Artificial Sequence <220> <223> Oligonucleoide Primer <400> 58 39 ctcgagaaaa gagttgttgg gggcacggat gcggatgag <210> 59 <211> 11 <212> PRT <213> Homo Sapien Phe Glu Val Phe Ser Gln Ser Ser Ser Leu Gly 5 <210> 60 <211> 32 <212> PRT <213> Homo Sapien Glu Ile Val Ala Pro Arg Glu Arg Ala Asp Arg Arg Gly Arg Lys Leu 5 10 Leu Cys Trp Arg Lys Pro Thr Lys Met Lys Gly Pro Arg Pro Ser His <210> 61 <211> 4933 <212> DNA <213> Homo Sapien <220> <221> CDS <222> (94)...(3222) <223> Nucleotide sequence encoding corin <300> <308> GenBank AF133845 <309> 1999-05-24 60 aaatcatccg tagtgcctcc ccgggggaca cgtagaggag agaaaagcga ccaagataaa 114 agtggacaga agaataagcg agacttttta tcc atg aaa cag tct cct gcc ctc Met Lys Gln Ser Pro Ala Leu 162 gct ccg gaa gag cgc tac cgc aga gcc ggg tcc cca aag ccg gtc ttg Ala Pro Glu Glu Arg Tyr Arg Arg Ala Gly Ser Pro Lys Pro Val Leu

aga gct gat gac aat aac atg ggc aat ggc tgc tct cag aag ctg gcg

210

## WO 01/57194 PCT/US01/03471 40/66

Arg	Ala 25	Asp	Asp	Asn	Asn	Met 30	Gly	Asn	Gly	Cys	Ser 35	Gln	Lys	Leu	Ala	
			ctc Leu													258
			gtt Val													306
			gtc Val 75													354
			atc Ile													402
			act Thr													450
			acg Thr													498
			tgt Cys													546
			acg Thr 155													594
			ttc Phe													642
			atc Ile													690
			ggc Gly													738
			gca Ala													786
			ccg Pro 235													834
			aat Asn													882
gga Gly	aag Lys 265	caa Gln	ttg Leu	ctc Leu	tgt Cys	gga Gly 270	agg Arg	ggt Gly	gag Glu	aac Asn	ttt Phe 275	ctg Leu	tgt Cys	gcc Ala	agt Ser	930

<b>WO</b> 01/57194		PCT/US01/03471				
	41/66					

								caa Gln								978
gac Asp	gac Asp	tgg Trp	agt Ser	gac Asp 300	gag Glu	gct Ala	cat His	tgc Cys	aac Asn 305	tgc Cys	agc Ser	gag Glu	aat Asn	ctg Leu 310	ttt Phe	1026
								aat Asn 320								1074
								gat Asp								1122
								gac Asp								1170
								tgt Cys								1218
								ctg Leu								1266
								gat Asp 400								1314
								gtc Val								1362
								ccc Pro								1410
								agt Ser								1458
			Thr		Glu		Cys	atg Met	Asn	Leu						1506
								agg Arg 480								1554
								gca Ala								1602
								acc Thr								1650
gtg	aat	aca	ggc	gag	cgt	atc	cct	cct	tgc	agg	gca	ttg	tgt	gaa	cac	1698

Val 520	Asn	Thr	Gly	Glu	Arg 525	Ile	Pro	Pro	Cys	Arg 530	Ala	Leu	Cys	Glu	His 535	
	aaa Lys															1746
	gaa Glu															1794
caa Gln	acc Thr	tgc Cys 570	ctg Leu	atg Met	cct Pro	gat Asp	gaa Glu 575	tat Tyr	gtg Val	gaa Glu	gaa Glu	tgc Cys 580	tca Ser	cct Pro	agt Ser	1842
	ttc Phe 585															1890
	ggc Gly															1938
	aaa Lys															1986
	cac His															2034
	gag Glu															2082
	cat His 665															2130
tgc Cys 680	tca Ser	gac Asp	agt Ser	tca Ser	gat Asp 685	gaa Glu	tgg Trp	gac Asp	tgt Cys	gtg Val 690	acc Thr	ctc Leu	tct Ser	ata Ile	aat Asn 695	2178
	aac Asn															2226
cat His	gtg Val	tgt Cys	gca Ala 715	gat Asp	ggc Gly	tgg Trp	cag Gln	gag Glu 720	ata Ile	ttg Leu	agt Ser	cag Gln	ctg Leu 725	gcc Ala	tgc Cys	2274
	cag Gln															2322
cag Gln	gag Glu 745	aaa Lys	gag Glu	ccg Pro	cgg Arg	tgg Trp 750	ctg Leu	aca Thr	tta Leu	cac His	tcc Ser 755	aac Asn	tgg Trp	gag Glu	agc Ser	2370
	aat Asn															2418

<b>WO</b> 01/57194	PCT/US01/03471

			agt Ser													2466
			gct Ala 795													2514
agt Ser	cgc Arg	cct Pro 810	gga Gly	agg Arg	tgg Trp	cca Pro	tgg Trp 815	cag Gln	tgt Cys	tct Ser	ctg Leu	cag Gln 820	agt Ser	gaa Glu	ccc Pro	2562
			atc Ile													2610
aca Thr 840			cac His													2658
gtg Val	gtg Val	ctt Leu	ggc Gly	atc Ile 860	aac Asn	aat Asn	cta Leu	gac Asp	cat His 865	cca Pro	tca Ser	gtg Val	ttc Phe	atg Met 870	cag Gln	2706
			gtg Val 875													2754
			tat Tyr													2802
			tac Tyr													2850
			gac Asp													2898
			cca Pro													2946
			tgt Cys 955													2994
			gct Ala													3042
		ĞĪy	gly aaa				Cys					ĞĪy				3090
	Phe		tta Leu			$\operatorname{Trp}$					Phe					3138
aaa	cct	ggc	gtt	tat	agt	aat	gtg	tca	tat	ttc	gtc	gaa	tgg	att	aaa	3186

Gly Pro Gly Val Ty	vr Ser Asn Val : 020	Ser Tyr Phe 1025		le Lys 030
aga cag att tac at Arg Gln Ile Tyr Il 1035	e Gln Thr Phe :		taa ttataagga *	3232
gatcagagac tittgcc cctgcagaga gctgtac aatttgcatg tittgtt cttatttcaa gttcaat ccaggcctaa gctaat tttaggctga ttaattt attcagtgt taacatt tttttattac acttcga tatacatgtg ccatgtt caagaataatt tacatgt tgagaactga aagttct ctggatgtat tacaaga attgaatgca aagttct ctggatgtat tatgggtttgtt atgggcata ttttagag ttaatatatt tttcagagaa tttaagac ggtttct gcactttac gcaactc aaaactta ggtttct gcactttac gcaactc aaaactta gacacta ttccaaatag atactac tctgtataaa ccaacac tctaatact gccacttac gcactttaca aaccac gcctttaaa caactac ttccaaatag atactac tctgtataaa cctaaga acatagttga aattcac tttagcctg ctcaact cttaatacaa cctaaga acatagtcaag acatagtcaag acataactac cctaaacac cctaacacac cctaaacac cctaacac ccac ccac cctaacac ccac cca	raga agcacttttc rtgg actaatttt rgaa agacttaca rtgc agcacaaaat rtgc agcacaaaat rtgg aatggtctct rtgga gacttgcttt rega gacttgctca reta cagtaattaa reta gagtatcaata reta cttatggga rtta cttgcgttca reta cttggagt reta cttcgaagt reta cttcataata reta catttata reta cttcataata reta cattttata reta cttcataata reta cattttata reta cattttat reta cattttat reta cattttat reta catttrat reta cattttat reta catttrat reta cattttat reta cattttat reta catttrat reta catttrat reta catttrat reta catttrat reta catttrat reta catttrat reta cattrat r	atggacagaa ttcaatttat aaagcaaaca tatcgactct tttatcctat ttgtttctca tctttttctt cgagcacaac cagtactcg tattacccc cagtgtcat ttaacacttg gaagagtctc attgagactg aatgaactgg aagaaatta tccataccaa aaaacatttt taaaatttta attagtggta acttcacacaca catgtatta cctccaccac cagtgttat ttaaatttta acttcatcac catgtattt tttctatact cagcaactga cagttgtagc	atgeteaate gtetttteacet teaageagaett teggegagattt aaa ageteagtge attetttaaee gtettttaaee gtettttaaee gtettteatetgatt teattegatt teettaatgtt teettaatgtt teettaaegt age acattattgg acattattgeate etaagaeetg teetaaagteata aagteattee etaaagteata agetaatttee gaaatgattt teaaagteata teagteagatt teaaaagtattt etaaaagtattt etaaaagtattt etaagteagg etgtatetttt etgategtea etgtatette etgategtea agetaatete etgategtea agetaatete etgategtea etgateatete etgategtea etgateatete attegtegtea agetaatetea attegtegtea etgateatetea attegtegtea etgateateatea attegtegtea etgateatea etgatea etgateatea etgateatea etgateatea etgateatea etgateatea etgateatea etgatea etgateatea etgateatea etgateatea etgateatea etgateatea etgatea etgateatea etgatea etgateatea etgatea etgate	gcactgca 3352 attittct 3412 tccttttg 3472 aatcaggt 3532 catagata 3592 agtggtaa 3652 acaattct 3712 tacatatg 3772 tacaaagc 3832 cataaattc 3892 cataaattc 38952 cataaattc 4012 ctcatagc 4072 ctatattt 4132 gcactatatt 4132 gcactatat 4252 ggtttaat 4252 ggtttaat 4312 aagctctag 4492 tcccacta 4492 tcccacta 4612 gcgtgtaa 4672 tcctcagca 4672 tcctcagca 4672 tccttcag 4732 attgtatg 4792 gtatcaac 4852
<213> Homo Sapien				
<pre>&lt;400&gt; 62 Met Lys Gln Ser Pr 1</pre>	5	10	1!	5
Gly Ser Pro Lys Pro 20		25	30	
Gly Cys Ser Gln Ly 35	s Leu Ala Thr . 40	Ala Asn Leu	Leu Arg Phe Lo	eu Leu
Leu Val Leu Ile Pr 50	55		60	
Leu Leu Ser Tyr Va 65	70	75		80
Gly Ser Glu Pro Le	5	90	9.	5
Ile Leu Thr Asn Th		105	110	
His Pro Asp Gln H	120		125	
Gly Asp Gln Ser H				
130 Ser Gln Cys Gln Me	135		140	

150 155 Ser Val Val Arg Asn Met Glu Met Glu Lys Phe Leu Lys Phe Phe Thr 165 170 Tyr Leu His Arg Leu Ser Cys Tyr Gln His Ile Met Leu Phe Gly Cys 185 180 Thr Leu Ala Phe Pro Glu Cys Ile Ile Asp Gly Asp Asp Ser His Gly 195 200 205 Leu Leu Pro Cys Arg Ser Phe Cys Glu Ala Ala Lys Glu Gly Cys Glu 215 220 Ser Val Leu Gly Met Val Asn Tyr Ser Trp Pro Asp Phe Leu Arg Cys 230 235 Ser Gln Phe Arg Asn Gln Thr Glu Ser Ser Asn Val Ser Arg Ile Cys 245 250 Phe Ser Pro Gln Glu Asn Gly Lys Gln Leu Leu Cys Gly Arg Gly 260 265 Glu Asn Phe Leu Cys Ala Ser Gly Ile Cys Ile Pro Gly Lys Leu Gln 275 280 285 Cys Asn Gly Tyr Asn Asp Cys Asp Asp Trp Ser Asp Glu Ala His Cys 295 300 Asn Cys Ser Glu Asn Leu Phe His Cys His Thr Gly Lys Cys Leu Asn 310 315 Tyr Ser Leu Val Cys Asp Gly Tyr Asp Asp Cys Gly Asp Leu Ser Asp 325 330 335 Glu Gln Asn Cys Asp Cys Asn Pro Thr Thr Glu His Arg Cys Gly Asp 345 340 Gly Arg Cys Ile Ala Met Glu Trp Val Cys Asp Gly Asp His Asp Cys 355 360 365 Val Asp Lys Ser Asp Glu Val Asn Cys Ser Cys His Ser Gln Gly Leu 370 375 380 Val Glu Cys Arg Asn Gly Gln Cys Ile Pro Ser Thr Phe Gln Cys Asp 390 395 Gly Asp Glu Asp Cys Lys Asp Gly Ser Asp Glu Glu Asn Cys Ser Val 410 415 405 Ile Gln Thr Ser Cys Gln Glu Gly Asp Gln Arg Cys Leu Tyr Asn Pro 425 430 Cys Leu Asp Ser Cys Gly Gly Ser Ser Leu Cys Asp Pro Asn Asn Ser 445 440 Leu Asn Asn Cys Ser Gln Cys Glu Pro Ile Thr Leu Glu Leu Cys Met 455 Asn Leu Pro Tyr Asn Ser Thr Ser Tyr Pro Asn Tyr Phe Gly His Arg 470 475 Thr Gln Lys Glu Ala Ser Ile Ser Trp Glu Ser Ser Leu Phe Pro Ala 485 490 Leu Val Gln Thr Asn Cys Tyr Lys Tyr Leu Met Phe Phe Ser Cys Thr 505 510 500 Ile Leu Val Pro Lys Cys Asp Val Asn Thr Gly Glu Arg Ile Pro Pro 515 520 525 Cys Arg Ala Leu Cys Glu His Ser Lys Glu Arg Cys Glu Ser Val Leu 540 535 Gly Ile Val Gly Leu Gln Trp Pro Glu Asp Thr Asp Cys Ser Gln Phe 550 555 Pro Glu Glu Asn Ser Asp Asn Gln Thr Cys Leu Met Pro Asp Glu Tyr 570 · 565 Val Glu Glu Cys Ser Pro Ser His Phe Lys Cys Arg Ser Gly Gln Cys 590 580 585 Val Leu Ala Ser Arg Arg Cys Asp Gly Gln Ala Asp Cys Asp Asp Asp 600 605 Ser Asp Glu Glu Asn Cys Gly Cys Lys Glu Arg Asp Leu Trp Glu Cys 615 620 Pro Ser Asn Lys Gln Cys Leu Lys His Thr Val Ile Cys Asp Gly Phe 635 630

Pro Asp Cys Pro Asp Tyr Met Asp Glu Lys Asn Cys Ser Phe Cys Gln

650 645 Asp Asp Glu Leu Glu Cys Ala Asn His Ala Cys Val Ser Arg Asp Leu 670 660 665 Trp Cys Asp Gly Glu Ala Asp Cys Ser Asp Ser Ser Asp Glu Trp Asp 675 680 685 Cys Val Thr Leu Ser Ile Asn Val Asn Ser Ser Ser Phe Leu Met Val 695 700 His Arg Ala Ala Thr Glu His His Val Cys Ala Asp Gly Trp Gln Glu 715 710 Ile Leu Ser Gln Leu Ala Cys Lys Gln Met Gly Leu Gly Glu Pro Ser 725 730 Val Thr Lys Leu Ile Gln Glu Gln Glu Lys Glu Pro Arg Trp Leu Thr 750 740 745 Leu His Ser Asn Trp Glu Ser Leu Asn Gly Thr Thr Leu His Glu Leu 755 760 Leu Val Asn Gly Gln Ser Cys Glu Ser Arg Ser Lys Ile Ser Leu Leu 770 775 780 Cys Thr Lys Gln Asp Cys Gly Arg Arg Pro Ala Ala Arg Met Asn Lys 790 795 Arg Ile Leu Gly Gly Arg Thr Ser Arg Pro Gly Arg Trp Pro Trp Gln 805 810 Cys Ser Leu Gln Ser Glu Pro Ser Gly His Ile Cys Gly Cys Val Leu 825 820 Ile Ala Lys Lys Trp Val Leu Thr Val Ala His Cys Phe Glu Gly Arg 835 840 845 Glu Asn Ala Ala Val Trp Lys Val Val Leu Gly Ile Asn Asn Leu Asp 855 860 His Pro Ser Val Phe Met Gln Thr Arg Phe Val Lys Thr Ile Ile Leu 870 875 His Pro Arg Tyr Ser Arg Ala Val Val Asp Tyr Asp Ile Ser Ile Val 885 890 895 Glu Leu Ser Glu Asp Ile Ser Glu Thr Gly Tyr Val Arg Pro Val Cys 905 Leu Pro Asn Pro Glu Gln Trp Leu Glu Pro Asp Thr Tyr Cys Tyr Ile 920 925 Thr Gly Trp Gly His Met Gly Asn Lys Met Pro Phe Lys Leu Gln Glu 935 940 Gly Glu Val Arg Ile Ile Ser Leu Glu His Cys Gln Ser Tyr Phe Asp 950 955 Met Lys Thr Ile Thr Thr Arg Met Ile Cys Ala Gly Tyr Glu Ser Gly 970 975 965 Thr Val Asp Ser Cys Met Gly Asp Ser Gly Gly Pro Leu Val Cys Glu 980 985 990 Lys Pro Gly Gly Arg Trp Thr Leu Phe Gly Leu Thr Ser Trp Gly Ser 995 1000 1005 Val Cys Phe Ser Lys Val Leu Gly Pro Gly Val Tyr Ser Asn Val Ser 1010 1015 1020 Tyr Phe Val Glu Trp Ile Lys Arg Gln Ile Tyr Ile Gln Thr Phe Leu 1025 1030 1035 Leu Asn

```
<210> 63
<211> 3696
<212> DNA
<213> Homo Sapien
```

<220> <221> CDS

<222> (41)...(3100)

<223> Nucleotide sequence encoding human entorkinase

<300> <308> GenBank HSU09860 <309> 1995-06-03 <400> 63 accagacagt tottaaatta gcaagcotto aaaaccaaaa atg ggg tog aaa aga 55 Met Gly Ser Lys Arg ggc ata tet tet agg cat cat tet etc age tec tat gaa atc atg ttt 103 Gly Ile Ser Ser Arg His His Ser Leu Ser Ser Tyr Glu Ile Met Phe gca gct ctc ttt gcc ata ttg gta gtg ctc tgt gct gga tta att gca 151 Ala Ala Leu Phe Ala Ile Leu Val Val Leu Cys Ala Gly Leu Ile Ala gta tcc tgc ctg aca atc aag gaa tcc caa cga ggt gca gca ctt gga Val Ser Cys Leu Thr Ile Lys Glu Ser Gln Arg Gly Ala Ala Leu Gly 199 cag agt cat gaa gcc aga gcg aca ttt aaa ata aca tcc gga gtt aca Gln Ser His Glu Ala Arg Ala Thr Phe Lys Ile Thr Ser Gly Val Thr 247 tat aat cct aat ttg caa gac aaa ctc tca gtg gat ttc aaa gtt ctt Tyr Asn Pro Asn Leu Gln Asp Lys Leu Ser Val Asp Phe Lys Val Leu 295 gct ttt gac ctt cag caa atg ata gat gag atc ttt cta tca agc aat 343 Ala Phe Asp Leu Gln Gln Met Ile Asp Glu Ile Phe Leu Ser Ser Asn 90 95 ctg aag aat gaa tat aag aac tca aga gtt tta caa ttt gaa aat ggc 391 Leu Lys Asn Glu Tyr Lys Asn Ser Arg Val Leu Gln Phe Glu Asn Gly age att ata gte gta ttt gae ett tte ttt gee eag tgg gtg tea gat 439 Ser Ile Ile Val Val Phe Asp Leu Phe Phe Ala Gln Trp Val Ser Asp 125 caa aat gta aaa gaa gaa ctg att caa ggc ctt gaa gca aat aaa tcc 487 Gln Asn Val Lys Glu Glu Leu Ile Gln Gly Leu Glu Ala Asn Lys Ser age caa ctg gtc act ttc cat att gat ttg aac age gtt gat atc cta 535 Ser Gln Leu Val Thr Phe His Ile Asp Leu Asn Ser Val Asp Ile Leu gac aag cta aca acc acc agt cat ctg gca act cca gga aat gtc tca Asp Lys Leu Thr Thr Ser His Leu Ala Thr Pro Gly Asn Val Ser 583 170 ata gag tgc ctg cct ggt tca agt cct tgt act gat gct cta acg tgt 631 Ile Glu Cys Leu Pro Gly Ser Ser Pro Cys Thr Asp Ala Leu Thr Cys 1.90 679 ata aaa gct gat tta ttt tgt gat gga gaa gta aac tgt cca gat ggt Ile Lys Ala Asp Leu Phe Cys Asp Gly Glu Val Asn Cys Pro Asp Gly 205 727 tct gac gaa gac aat aaa atg tgt gcc aca gtt tgt gat gga aga ttt

## WO 01/57194 PCT/US01/03471 48/66

Ser	Asp 215	Glu	Asp	Asn	Lys	Met 220	Cys	Ala	Thr	Val	Cys 225	Asp	Gly	Arg	Phe	
ttg Leu 230	tta Leu	act Thr	gga Gly	tca Ser	tct Ser 235	gl <sup>à</sup> aaa	tct Ser	ttc Phe	cag Gln	gct Ala 240	act Thr	cat His	tat Tyr	cca Pro	aaa Lys 245	775
											ata Ile					823
											aat Asn					871
											agc Ser					919
											aga Arg 305					967
											gaa Glu					1015
ggc gly	ttt Phe	aat Asn	gca Ala	aca Thr 330	tat Tyr	act Thr	gca Ala	ttt Phe	aac Asn 335	agc Ser	agt Ser	gag Glu	ctt Leu	aat Asn 340	aat Asn	1063
											ttt Phe					1111
											att Ile					1159
											act Thr 385					1207
											gga Gly					1255
											act Thr					1303
tgc Cys	ctt Leu	agt Ser	ttc Phe 425	tgg Trp	tat Tyr	cat His	atg Met	tat Tyr 430	ggt Gly	gaa Glu	aat Asn	gtc Val	cat His 435	aaa Lys	tta Leu	1351
											aag Lys					1399
											gga Gly 465					1447

aat Asn 470	gaa Glu	aca Thr	gtt Val	·aaa Lys	ttt Phe 475	aag Lys	gtt Val	gct Ala	ttt Phe	aat Asn 480	gct Ala	ttt Phe	aaa Lys	aac Asn	aag Lys 485		1495
											cta Leu					:	1543
											gtg Val						1591
											gag Glu					,	1639
											agc Ser 545						1687
											gga Gly						1735
											aac Asn						1783
											tta Leu						1831
											acc Thr						1879
											gga Gly 625						1927
											gag Glu						1975
gac Asp	cat His	ttt Phe	caa Gln	tgt Cys 650	aaa Lys	aat Asn	gga Gly	gag Glu	tgt Cys 655	gtt Val	cca Pro	ctg Leu	gtg Val	aat Asn 660	ctc Leu		2023
											gat Asp						2071
											ggt Gly						2119
											gag Glu 705						2167
cag	att	tca	aat	gat	gtt	tgt	caa	ctg	ctg	gga	cta	<b>a</b> aa	agt	gga	aac		2215

Gln 710	Ile	Ser	Asn	Asp	Val 715	Cys	Gln	Leu	Leu	Gly 720	Leu	Gly	Ser	Gly	Asn 725		
										gga Gly						2263	3
										aca Thr						231	1
										aac Asn						2359	9
										aag Lys						240	7
										gtg Val 800						245!	5
										agc Ser						250	3
										tta Leu						255	1
										aat Asn						259:	9
										ata Ile						264'	7
										atg Met 880						269!	5
										tgt Cys						274	3
										att Ile						279:	1
										ttg Leu						283	9
										cag Gln						288	7
att Ile 950	act Thr	gaa Glu	aat Asn	atg Met	ata Ile 955	tgt Cys	gca Ala	ggc	tat Tyr	gaa Glu 960	gaa Glu	gga Gly	gga Gly	ata Ile	gat Asp 965	293	5

														aac Asn 980		2983
				Ala					Phe					gcc Ala		3031
			Pro					Arg					Thr	gaa Glu		3079
		Ser	ttt Phe			tag *	cgca	attt	ctt a	aact	caaad	ca gg	gaaag	gtego	C	3130
aaaaaaacaactaaacaacaacaacaacaacaacaacaac	agtta aaatt ataca aatga acact catgt ctag	acc act act att att act acca acca acca	aaagg aaaaa agtat atttt agaaa ctgtt aaact	gtttt ataat cttatt attto cggaa ccaat	et at ca at ca	tctt atto gtgaa ctact cagaa aact cttct	cacct cacca gette attta cgcca cctgt	tate  g tate  g tate  g ttate  a aaa  a taa  c ctt	gtcaa gcaat sttct saaag aaaga aattt sttct	atga caca ctca ggga aaag ctct	aato gaat caga tgtt aaaa agtt	getag taact atcto tattt ataas tccag atttt	ggg gett a cat i	ggcca aaaat cttta aagca gtttt cagtt acata	atttta agggaa caccat aaaatt atatac ctcca ctgctg atgcat ggcctg	3190 3250 3310 3370 3430 3490 3550 3610 3670 3696
<213	0> 64 L> 10 2> PE 3> Ho	)19 RT	Sapie	en												
			_													
<400 Met	)> 64	1	_	Arg	Gly	Ile	Ser	Ser	_	His	His	Ser	Leu	Ser	Ser	
<400 Met 1	0> 64 Gly	1 Ser	Lys	Arg 5	_				10	_		_	_	Ser 15 Leu		
<400 Met 1 Tyr	)> 64 Gly Glu	Ser	Lys Met 20	Arg 5 Phe	Ala	Ala	Leu	Phe 25	10 Ala	Ile	Leu	Val	Val 30	15	Cys	
<400 Met 1 Tyr	O> 64 Gly Glu Gly	Ser Ile Leu 35	Lys Met 20 Ile	Arg 5 Phe Ala	Ala Val	Ala Ser	Leu Cys 40	Phe 25 Leu	10 Ala Thr	Ile Ile	Leu Lys	Val Glu 45	Val 30 Ser	15 Leu	Cys Arg	
<400 Met 1 Tyr Ala Gly	Gly Glu Gly Gly Ala	Ser Ile Leu 35 Ala	Lys Met 20 Ile Leu	Arg 5 Phe Ala Gly	Ala Val Gln	Ala Ser Ser 55	Leu Cys 40 His	Phe 25 Leu Glu	10 Ala Thr Ala	Ile Ile Arg	Leu Lys Ala 60	Val Glu 45 Thr	Val 30 Ser Phe	15 Leu Gln Lys	Cys Arg	
<400 Met 1 Tyr Ala Gly Thr	Gly Glu Gly Ala 50 Ser	Ser Ile Leu 35 Ala	Lys Met 20 Ile Leu Val	Arg 5 Phe Ala Gly Thr Leu	Ala Val Gln Tyr 70	Ala Ser Ser 55 Asn	Leu Cys 40 His Pro	Phe 25 Leu Glu Asn	10 Ala Thr Ala Leu Gln	Ile Ile Arg Gln 75	Leu Lys Ala 60 Asp	Val Glu 45 Thr Lys	Val 30 Ser Phe Leu	15 Leu Gln Lys Ser Glu	Cys Arg Ile Val 80	
<400 Met 1 Tyr Ala Gly Thr 65 Asp	Gly Gly Gly Ala 50 Ser	Ser Ile Leu 35 Ala Gly Lys Ser	Lys Met 20 Ile Leu Val Val Ser	Arg 5 Phe Ala Gly Thr Leu 85 Asn	Ala Val Gln Tyr 70 Ala Leu	Ala Ser Ser 55 Asn Phe Lys	Leu Cys 40 His Pro Asp	Phe 25 Leu Glu Asn Leu Glu	10 Ala Thr Ala Leu Gln 90 Tyr	Ile Ile Arg Gln 75 Gln	Leu Lys Ala 60 Asp Met	Val Glu 45 Thr Lys Ile	Val 30 Ser Phe Leu Asp	15 Leu Gln Lys Ser Glu 95 Val	Cys Arg Ile Val 80 Ile	
<400 Met 1 Tyr Ala Gly Thr 65 Asp	Gly Gly Gly Ala 50 Ser Phe Leu	Ser Ile Leu 35 Ala Gly Lys Ser Glu	Lys Met 20 Ile Leu Val Val Ser	Arg 5 Phe Ala Gly Thr Leu 85 Asn	Ala Val Gln Tyr 70 Ala Leu	Ala Ser Ser 55 Asn Phe Lys	Leu Cys 40 His Pro Asp Asn Ile	Phe 25 Leu Glu Asn Leu Glu 105	10 Ala Thr Ala Leu Gln 90 Tyr	Ile Ile Arg Gln 75 Gln Lys	Leu Lys Ala 60 Asp Met	Val Glu 45 Thr Lys Ile Ser Leu	Val 30 Ser Phe Leu Asp Arg	15 Leu Gln Lys Ser Glu 95 Val	Cys Arg Ile Val 80 Ile Leu	
<400 Met 1 Tyr Ala Gly Thr 65 Asp Phe	Gly Gly Gly Ala 50 Ser Phe Leu Phe	Ser Ile Leu 35 Ala Gly Lys Ser Glu 115	Lys Met 20 Ile Leu Val Val Ser 100 Asn	Arg 5 Phe Ala Gly Thr Leu 85 Asn	Ala Val Gln Tyr 70 Ala Leu Ser	Ala Ser Ser 55 Asn Phe Lys Ile Asn	Leu Cys 40 His Pro Asp Asn Ile 120	Phe 25 Leu Glu Asn Leu Glu 105 Val	10 Ala Thr Ala Leu Gln 90 Tyr Val	Ile Ile Arg Gln 75 Gln Lys Phe	Leu Lys Ala 60 Asp Met Asn Asp	Val Glu 45 Thr Lys Ile Ser Leu 125	Val 30 Ser Phe Leu Asp Arg 110 Phe	15 Leu Gln Lys Ser Glu 95 Val	Cys Arg Ile Val 80 Ile Leu Ala	
<400 Met 1 Tyr Ala Gly Thr 65 Asp Phe Gln Gln	Gly Gly Ala 50 Ser Phe Leu Phe Trp 130	Ser Leu 35 Ala Gly Lys Ser Glu 115 Val	Lys Met 20 Ile Leu Val Val Ser 100 Asn	Arg 5 Phe Ala Gly Thr Leu 85 Asn Gly Asp	Ala Val Gln Tyr 70 Ala Leu Ser Gln Ser	Ala Ser Ser 55 Asn Phe Lys Ile Asn 135	Leu Cys 40 His Pro Asp Asn Ile 120 Val	Phe 25 Leu Glu Asn Leu Glu 105 Val	Thr Ala Leu Gln 90 Tyr Val	Ile Ile Arg Gln 75 Gln Lys Phe Glu Phe	Leu Lys Ala 60 Asp Met Asn Asp	Val Glu 45 Thr Lys Ile Ser Leu 125 Ile	Val 30 Ser Phe Leu Asp Arg 110 Phe	15 Leu Gln Lys Ser Glu 95 Val Phe	Cys Arg Ile Val 80 Ile Leu Ala Leu Asn	
<400 Met 1 Tyr Ala Gly Thr 65 Asp Phe Gln Gln Glu 145	Gly Gly Gly Ala 50 Ser Phe Leu Phe Trp 130 Ala	Ser Ile Leu 35 Ala Gly Lys Ser Glu 115 Val	Lys Met 20 Ile Leu Val Val Ser 100 Asn Ser Lys	Arg 5 Phe Ala Gly Thr Leu 85 Asn Gly Asp Ser Leu	Ala Val Gln Tyr 70 Ala Leu Ser Gln Ser 150	Ala Ser Ser 55 Asn Phe Lys Ile Asn 135 Gln	Leu Cys 40 His Pro Asp Asn Ile 120 Val Leu	Phe 25 Leu Glu Asn Leu Glu 105 Val Lys	10 Ala Thr Ala Leu Gln 90 Tyr Val Glu Thr	Ile Ile Arg Gln 75 Gln Lys Phe Glu Phe 155	Leu Lys Ala 60 Asp Met Asn Asp Leu 140 His	Val Glu 45 Thr Lys Ile Ser Leu 125 Ile Ile	Val 30 Ser Phe Leu Asp 110 Phe Gln	15 Leu Gln Lys Ser Glu 95 Val Phe Gly Leu	Cys Arg Ile Val 80 Ile Leu Ala Leu Asn 160	
<400 Met 1 Tyr Ala Gly Thr 65 Asp Phe Gln Gln Glu 145 Ser	Gly Gly Gly Ala 50 Ser Phe Leu Phe Trp 130 Ala Val	Ser Ile Leu 35 Ala Gly Lys Ser Glu 115 Val Asn Asp	Lys Met 20 Ile Leu Val Val Ser 100 Asn Ser Lys Ile Val	Arg 5 Phe Ala Gly Thr Leu 85 Asn Gly Asp Ser Leu 165	Ala Val Gln Tyr 70 Ala Leu Ser Gln Ser 150 Asp	Ala Ser Ser 55 Asn Phe Lys Ile Asn 135 Gln Lys	Leu Cys 40 His Pro Asp Asn Ile 120 Val Leu Leu	Phe 25 Leu Glu Asn Leu Glu 105 Val Lys Val Thr	Thr Ala Leu Gln 90 Tyr Val Glu Thr	Ile Ile Arg Gln 75 Gln Lys Phe Glu Phe 155 Thr	Leu Lys Ala 60 Asp Met Asn Asp Leu 140 His	Val Glu 45 Thr Lys Ile Ser Leu 125 Ile Ile His	Val 30 Ser Phe Leu Asp 110 Phe Gln Asp Leu Pro	15 Leu Gln Lys Ser Glu 95 Val Phe Gly Leu	Cys Arg Ile Val 80 Ile Leu Ala Leu Asn 160 Thr	
<400 Met 1 Tyr Ala Gly Thr 65 Asp Phe Gln Gln Glu 145 Ser Pro	Gly Gly Gly Ala 50 Ser Phe Leu Phe Trp 130 Ala Val	Ser Ile Leu 35 Ala Gly Lys Ser Glu 115 Val Asn Asp	Lys Met 20 Ile Leu Val Val Ser 100 Asn Ser Lys Ile Val 180	Arg 5 Phe Ala Gly Thr Leu 85 Asn Gly Asp Ser Leu 165 Ser	Ala Val Gln Tyr 70 Ala Leu Ser Gln Ser 150 Asp	Ala Ser Ser 55 Asn Phe Lys Ile Asn 135 Gln Lys Glu	Leu Cys 40 His Pro Asp Asn Ile 120 Val Leu Leu Cys	Phe 25 Leu Glu Asn Leu 105 Val Lys Val Thr Leu 185	10 Ala Thr Ala Leu Gln 90 Tyr Val Glu Thr Thr 170 Pro	Ile Ile Arg Gln 75 Gln Lys Phe Glu Phe 155 Thr Gly	Leu Lys Ala 60 Asp Met Asn Asp Leu 140 His Ser Ser	Val Glu 45 Thr Lys Ile Ser Leu 125 Ile Ile His Ser	Val 30 Ser Phe Leu Asp Arg 110 Phe Gln Asp Leu Pro	15 Leu Gln Lys Ser Glu 95 Val Phe Gly Leu Ala 175	Cys Arg Ile Val 80 Ile Leu Ala Leu Asn 160 Thr	

Cys 225	Asp	Gly	Arg	Phe	Leu 230	Leu	Thr	Gly	Ser	Ser 235	Gly	Ser	Phe	Gln	Ala 240
	His	Tyr	Pro			Ser	Glu	Thr			Val	Cys	Gln	Trp 255	
Ile	Arg	Val	Asn 260	245 Gln	Gly	Leu	Ser	Ile 265	250 Lys	Leu	Ser	Phe	Asp 270		Phe
Asn	Thr	Tyr 275	Tyr	Thr	Aśp	Ile	Leu 280		Ile	Tyr	Glu	Gly 285		Gly	Ser
Ser	Lys 290		Leu	Arg	Ala	Ser 295		Trp	Glu	Thr	Asn 300		Gly	Thr	Ile
Arg 305	Ile	Phe	Ser	Asn	Gln 310	Val	Thr	Ala	Thr	Phe 315	Leu	Ile	Glu	Ser	Asp 320
	Ser	Asp	Tyr	Val 325		Phe	Asn	Ala	Thr 330	Tyr	Thr	Ala	Phe	Asn 335	Ser
Ser	Glu	Leu	Asn 340		Tyr	Glu	Lys	Ile 345	Asn	Cys	Asn	Phe	Glu 350	Asp	Gly
Phe	Cys	Phe 355	Trp	Val	Gln	Asp	Leu 360	Asn	Asp	Asp	Asn	Glu 365	Trp	Glu	Arg
Ile	Gln 370	Gly	Ser	Thr	Phe	Ser 375	Pro	Phe	Thr	Gly	Pro 380	Asn	Phe	Asp	His
Thr 385	Phe	Gly	Asn	Ala	Ser 390	Gly	Phe	Tyr	Ile	Ser 395	Thr	Pro	Thr	Gly	Pro 400
Gly	Gly	Arg	Gln	Glu 405	Arg	Val	Gly	Leu	Leu 410	Ser	Leu	Pro	Leu	Asp 415	Pro
Thr	Leu	Glu	Pro 420	Ala	Cys	Leu	Ser	Phe 425	Trp	Tyr	His	Met	Tyr 430	Gly	Glu
Asn	Val	His 435	Lys	Leu	Ser	Ile	Asn 440	Ile	Ser	Asn	Asp	Gln 445	Asn	Met	Glu
Lys	Thr 450	Val	Phe	Gln	Lys	Glu 455	Gly	Asn	Tyr	Gly	Asp 460	Asn	Trp	Asn	Tyr
465			Thr		470					475					480
			Asn	485					490					495	
Leu	Thr	Tyr	Gly 500	Ile	Cys	Asn	Gly	Ser 505	Leu	Tyr	Pro	Glu	Pro 510	Thr	Leu
		515	Pro				520					525			
	530	_	Glu			535					540				
545			Asn		550					555					560
_	_		Ile	565					570					575	
	_		Val 580					585					590		
		595	Tyr				600					605			
	610	_				615					620		_	_	Arg
625	-		Lys		630					635					640
			Lys	645					650			_		655	
			Asn 660		-	_	_	665			-		670	_	
_		675	Asp	_		_	680					685			
	690		Arg			695					700				
Glu 705	Asn	Trp	Thr	Thr	Gln 710	Ile	Ser	Asn	Asp	Val 715	Cys	Gln	Leu	Leu	Gly 720

```
Leu Gly Ser Gly Asn Ser Ser Lys Pro Ile Phe Ser Thr Asp Gly Gly
                                       730
                                                             735
                 725
Pro Phe Val Lys Leu Asn Thr Ala Pro Asp Gly His Leu Ile Leu Thr
             740
                                   745
Pro Ser Gln Gln Cys Leu Gln Asp Ser Leu Ile Arg Leu Gln Cys Asn
                              760
                                                    765
His Lys Ser Cys Gly Lys Lys Leu Ala Ala Gln Asp Ile Thr Pro Lys
                          775
Ile Val Gly Gly Ser Asn Ala Lys Glu Gly Ala Trp Pro Trp Val Val
                     790
                                           795
Gly Leu Tyr Tyr Gly Gly Arg Leu Leu Cys Gly Ala Ser Leu Val Ser
                 805
                                       810
                                                             815
Ser Asp Trp Leu Val Ser Ala Ala His Cys Val Tyr Gly Arg Asn Leu
            820
                                   825
                                                        830
Glu Pro Ser Lys Trp Thr Ala Ile Leu Gly Leu His Met Lys Ser Asn
                              840
                                                    845
Leu Thr Ser Pro Gln Thr Val Pro Arg Leu Ile Asp Glu Ile Val Ile
                          855
Asn Pro His Tyr Asn Arg Arg Lys Asp Asn Asp Ile Ala Met Met
                     870
                                           875
His Leu Glu Phe Lys Val Asn Tyr Thr Asp Tyr Ile Gln Pro Ile Cys
                 885
                                       890
                                                             895
Leu Pro Glu Glu Asn Gln Val Phe Pro Pro Gly Arg Asn Cys Ser Ile
                                                        910
                                   905
            900
Ala Gly Trp Gly Thr Val Val Tyr Gln Gly Thr Thr Ala Asn Ile Leu
        915
                              920
                                                    925
Gln Glu Ala Asp Val Pro Leu Leu Ser Asn Glu Arg Cys Gln Gln Gln
                          935
                                                940
Met Pro Glu Tyr Asn Ile Thr Glu Asn Met Ile Cys Ala Gly Tyr Glu
                      950
                                           955
Glu Gly Gly Ile Asp Ser Cys Gln Gly Asp Ser Gly Pro Leu Met
                                       970
                 965
Cys Gln Glu Asn Asn Arg Trp Phe Leu Ala Gly Val Thr Ser Phe Gly
                                  985
            980
Tyr Lys Cys Ala Leu Pro Asn Arg Pro Gly Val Tyr Ala Arg Val Ser
        995
                              1000
                                                    1005
Arg Phe Thr Glu Trp Ile Gln Ser Phe Leu His
    1010
                          1015
<210> 65
<211> 1500
<212> DNA
<213> Homo Sapien
<220>
<221> CDS
<222> (62)...(1318)
<223> Nucleotide sequence encoding human airway
      trypsin-like protease
<300>
<308> GenBank AB002134
<309> 1998-06-04
<400> 65
gagtgggaat ctcaaagcag ttgagtaggc agaaaaaaga acctcttcat taaggattaa
                                                                            60
a atg tat agg cca gca cgt gta act tcg act tca aga ttt ctg aat cca
Met Tyr Arg Pro Ala Arg Val Thr Ser Thr Ser Arg Phe Leu Asn Pro
                                                                           109
                    5
tat gta gta tgt ttc att gtc gtc gca ggg gta gtg atc ctg gca gtc Tyr Val Val Cys Phe Ile Val Val Ala Gly Val Val Ile Leu Ala Val
```

			20				25			30			
											tct Ser		205
											agt Ser		253
											aga Arg		301
											aat Asn 95		349
											ggt Gly		397
_	_ ~	_	_	_	_				_		aat Asn		445
											ctg Leu	•	493
											tca Ser		541
											gcc Ala 175		589
											act Thr		637
											aat Asn		685
											ctg Leu		733
											att Ile		781
											gta Val 255		829
											aat Asn		877

	gca Ala															925
	agt Ser 290															973
act Thr 305	gct Ala	tat Tyr	gta Val	aca Thr	gga Gly 310	tgg Trp	ggc Gly	gct Ala	caa Gln	gaa Glu 315	tat Tyr	gct Ala	ggc	cac His	aca Thr 320	1021
gtt Val	cca Pro	gag Glu	cta Leu	agg Arg 325	caa Gln	gga Gly	cag Gln	gtc Val	aga Arg 330	ata Ile	ata Ile	agt Ser	aat Asn	gat Asp 335	gta Val	1069
	aat Asn															1117
tgt Cys	gct Ala	gga Gly 355	gta Val	cct Pro	caa Gln	ggt Gly	gga Gly 360	gtg Val	gac Asp	gca Ala	tgt Cys	cag Gln 365	ggt Gly	gac Asp	tct Ser	1165
	ggc Gly 370															1213
382 Gly aaa	ata Ile	gta Val	agc Ser	tgg Trp	gga Gly 390	gat Asp	cag Gln	tgt Cys	ggc Gly	ctg Leu 395	ccg Pro	gat Asp	aag Lys	cca Pro	gga Gly 400	1261
	tat Tyr															1309
	atc Ile	_	tgca	aacaa	agt g	gcate	cccts	gt to	gcaaa	agtc	t gta	atgc	aggt	¢		1358
aat		cat o	cttgt	taca	at aa	atat									tgtcct ttatta	1418 1478 1500
<21:	0> 66 1> 43 2> PI 3> Ho	L8 RT	Sapie	en											*	•
	0> 66 Tyr		Pro	Ala	Arg	Val	Thr	Ser	Thr	Ser	Arg	Phe	Leu	Asn	Pro	
1 Tyr	Val	Val	Cys	5 Phe	Ile	Val	Val		10 Gly	Val	Val	Ile		15 Ala	Val	
Thr	Ile	Ala 35	∠∪ Leu	Leu	Val	Tyr	Phe 40	25 Leu	Ala	Phe	Asp	Gln 45	Lys Lys	Ser	Tyr	
Phe	Tyr 50		Ser	Ser	Phe	Gln 55		Leu	Asn	Val	Glu 60		Asn	Ser	Gln	
Leu 65	Asn	Ser	Pro	Ala	Thr 70		Glu	Tyr	Arg	Thr 75		Ser	Gly	Arg	Ile 80	
	Ser	Leu	Ile	Thr 85	Lys	Thr	Phe	Lys	Glu 90	Ser	Asn	Leu	Arg	Asn 95	Gln	

```
Phe Ile Arg Ala His Val Ala Lys Leu Arg Gln Asp Gly Ser Gly Val
            100
                                105
Arg Ala Asp Val Val Met Lys Phe Gln Phe Thr Arg Asn Asn Asn Gly
                            120
        115
                                                125
Ala Ser Met Lys Ser Arg Ile Glu Ser Val Leu Arg Gln Met Leu Asn
                        135
                                            140
Asn Ser Gly Asn Leu Glu Ile Asn Pro Ser Thr Glu Ile Thr Ser Leu
                    150
                                        155
Thr Asp Gln Ala Ala Ala Asn Trp Leu Ile Asn Glu Cys Gly Ala Gly
                                    170
Pro Asp Leu Ile Thr Leu Ser Glu Gln Arg Ile Leu Gly Gly Thr Glu
                                185
                                                    190
            180
Ala Glu Glu Gly Ser Trp Pro Trp Gln Val Ser Leu Arg Leu Asn Asn
                            200
                                                205
        195
Ala His His Cys Gly Gly Ser Leu Ile Asn Asn Met Trp Ile Leu Thr
                        215
                                            220
Ala Ala His Cys Phe Arg Ser Asn Ser Asn Pro Arg Asp Trp Ile Ala
                    230
                                        235
Thr Ser Gly Ile Ser Thr Thr Phe Pro Lys Leu Arg Met Arg Val Arg
                                    250
Asn Ile Leu Ile His Asn Asn Tyr Lys Ser Ala Thr His Glu Asn Asp
            260
                                265
Ile Ala Leu Val Arg Leu Glu Asn Ser Val Thr Phe Thr Lys Asp Ile
                            280
                                                285
        275
His Ser Val Cys Leu Pro Ala Ala Thr Gln Asn Ile Pro Pro Gly Ser
                        295
                                            300
Thr Ala Tyr Val Thr Gly Trp Gly Ala Gln Glu Tyr Ala Gly His Thr
                                                             320
                    310
                                        315
Val Pro Glu Leu Arg Gln Gly Gln Val Arg Ile Ile Ser Asn Asp Val
                325
                                    330
                                                         335
Cys Asn Ala Pro His Ser Tyr Asn Gly Ala Ile Leu Ser Gly Met Leu
            340
                                345
                                                     350
Cys Ala Gly Val Pro Gln Gly Gly Val Asp Ala Cys Gln Gly Asp Ser
        355
                            360
                                                 365
Gly Gly Pro Leu Val Gln Glu Asp Ser Arg Arg Leu Trp Phe Ile Val
                        375
                                            380
Gly Ile Val Ser Trp Gly Asp Gln Cys Gly Leu Pro Asp Lys Pro Gly
                    390
                                        395
Val Tyr Thr Arg Val Thr Ala Tyr Leu Asp Trp Ile Arg Gln Gln Thr
                405
                                    410
Gly Ile
<210> 67
<211> 1783
<212> DNA
<213> Homo Sapien
<220>
<221> CDS
<222> (246)...(1499)
<223> Nucleotide sequence encoding human hepsin
<308> GenBank M18930
<309> 1993-06-11
<400> 67
tegagecege tttecaggga cectacetga gggeceaeag gtgaggeage etggeetage
aggececacg ceaeegeete tgeeteeagg eegeeegetg etgeggggee accatgetee
tgcccaggcc tggagactga cccgaccccg gcactacctc gaggctccgc ccccacctgc
tggaccccag ggtcccaccc tggcccagga ggtcagccag ggaatcatta acaagaggca
```

60

120

180

240

gtgac		t Al								ır Va					cc aga er Arg 15	1
ccc a Pro L																338
atc g Ile G																386
cag g Gln G																434
atg g Met V																482
cgc t Arg S 80																530
ctc a Leu A																578
aat g Asn G	gly gc	acg Thr	tcg Ser 115	gly ggc	ttc Phe	ttc Phe	tgt Cys	gtg Val 120	gac Asp	gag Glu	elà aaa	agg Arg	ctg Leu 125	ccc Pro	cac His	626
acc c Thr G	ln															674
cgt t Arg P 1																722
gtg g Val A 160																770
tgg c Trp G																818
ctg c Leu L	etc Leu	tcc Ser	999 195	gac Asp	tgg Trp	gtg Val	ctg Leu	aca Thr 200	gcc Ala	gcc Ala	cac His	tgc Cys	ttc Phe 205	ccg Pro	gag Glu	866
cgg a Arg A	Asn	cgg Arg 210	gtc Val	ctg Leu	tcc Ser	cga Arg	tgg Trp 215	cga Arg	gtg Val	ttt Phe	gcc Ala	ggt Gly 220	gcc Ala	gtg Val	gcc Ala	914
cag g Gln A 2	gcc Ala 225	tct Ser	ccc Pro	cac His	ggt Gly	ctg Leu 230	cag Gln	ctg Leu	glà aaa	gtg Val	cag Gln 235	gct Ala	gtg Val	gtc Val	tac Tyr	962
cac g His G	ily igg	ggc Gly	tat Tyr	ctt Leu	ccc Pro	ttt Phe	cgg Arg	gac Asp	ccc Pro	aac Asn	agc Ser	gag Glu	gag Glu	aac Asn	agc Ser	1010

WO 01/57194		PCT/US01/03471
	58/66	

240	245	250	255					
	Val His Leu Ser	agt ccc ctg ccc ctc Ser Pro Leu Pro Leu 265						
		gcc ggc cag gcc ctg Ala Gly Gln Ala Leu 285						
		ggc aac acg cag tac Gly Asn Thr Gln Tyr 300						
		cga gtc ccc ata atc Arg Val Pro Ile Ile 315						
		gga aac cag atc aag Gly Asn Gln Ile Lys 330						
atg ttc tgt gct ggc Met Phe Cys Ala Gly 340	Tyr Pro Glu Gly	ggc att gat gcc tgc Gly Ile Asp Ala Cys 345	cag ggc 1298 Gln Gly 350					
		gac agc atc tct cgg Asp Ser Ile Ser Arg 365						
cgt tgg cgg ctg tgt Arg Trp Arg Leu Cys 370	ggc att gtg agt Gly Ile Val Ser 375	tgg ggc act ggc tgt Trp Gly Thr Gly Cys 380	gcc ctg 1394 Ala Leu					
		gtc agt gac ttc cgg Val Ser Asp Phe Arg 395						
		gaa gcc agc ggc atg Glu Ala Ser Gly Met 410						
cag ctc tga ccggtgg Gln Leu *	ctt ctcgctgcgc ag	cctccagg gcccgaggtg	1539					
atcccggtgg tgggatccac gctgggccga ggatgggacg tttttcttct tgggcccggt ccacaggtcc aaggacaccc tccctccagg gtcctctctt ccacagtggc gggcccactc agccccgaga ccacccaacc tcaccctcct gacccccatg taaatattgt tctgctgtct gggactcctg tctaggtgcc cctgatgatg ggatgctctt taaataataa agatggtttt gatt								
<210> 68 <211> 417 <212> PRT <213> Homo Sapien								
		Val Pro Cys Cys Ser						
1 5 Lys Val Ala Ala Leu 20		10 Leu Leu Leu Leu Thr 30	15 Ala Ile					

```
Gly Ala Ala Ser Trp Ala Ile Val Ala Val Leu Leu Arg Ser Asp Gln
                          40
Glu Pro Leu Tyr Pro Val Gln Val Ser Ser Ala Asp Ala Arg Leu Met
                      55
Val Phe Asp Lys Thr Glu Gly Thr Trp Arg Leu Leu Cys Ser Ser Arg
                  70
                                   75
Ser Asn Ala Arg Val Ala Gly Leu Ser Cys Glu Glu Met Gly Phe Leu
              85
                                 90
Arg Ala Leu Thr His Ser Glu Leu Asp Val Arg Thr Ala Gly Ala Asn
           100
                              105
                                                 110
Gly Thr Ser Gly Phe Phe Cys Val Asp Glu Gly Arg Leu Pro His Thr
                          120
       115
                                             125
Gln Arg Leu Leu Glu Val Ile Ser Val Cys Asp Cys Pro Arg Gly Arg
                    135
                                        140
Phe Leu Ala Ala Ile Cys Gln Asp Cys Gly Arg Arg Lys Leu Pro Val
                                     155
                 150
Asp Arg Ile Val Gly Gly Arg Asp Thr Ser Leu Gly Arg Trp Pro Trp
               165
                                  170
                                                     175
Gln Val Ser Leu Arg Tyr Asp Gly Ala His Leu Cys Gly Gly Ser Leu
           180
                             185
                                                 190
Leu Ser Gly Asp Trp Val Leu Thr Ala Ala His Cys Phe Pro Glu Arg
       195
                          200
                                             205
Asn Arg Val Leu Ser Arg Trp Arg Val Phe Ala Gly Ala Val Ala Gln
           215
                               220
Ala Ser Pro His Gly Leu Gln Leu Gly Val Gln Ala Val Val Tyr His
                   230
                                      235
Gly Gly Tyr Leu Pro Phe Arg Asp Pro Asn Ser Glu Glu Asn Ser Asn
              245
                                250
Asp Ile Ala Leu Val His Leu Ser Ser Pro Leu Pro Leu Thr Glu Tyr
                              265
                                                 270
           260
Ile Gln Pro Val Cys Leu Pro Ala Ala Gly Gln Ala Leu Val Asp Gly
       275
                         280
                                             285
Lys Ile Cys Thr Val Thr Gly Trp Gly Asn Thr Gln Tyr Tyr Gly Gln
                      295
                                          300
Gln Ala Gly Val Leu Gln Glu Ala Arg Val Pro Ile Ile Ser Asn Asp
                   310
                                      315
                                                         320
Val Cys Asn Gly Ala Asp Phe Tyr Gly Asn Gln Ile Lys Pro Lys Met
                                  330
               325
Phe Cys Ala Gly Tyr Pro Glu Gly Gly Ile Asp Ala Cys Gln Gly Asp
           340
                              345
                                                 350
Ser Gly Gly Pro Phe Val Cys Glu Asp Ser Ile Ser Arg Thr Pro Arg
       355
                          360
                                             365
Trp Arg Leu Cys Gly Ile Val Ser Trp Gly Thr Gly Cys Ala Leu Ala
                   375
                                      380
Gln Lys Pro Gly Val Tyr Thr Lys Val Ser Asp Phe Arg Glu Trp Ile
                   390
                                      395
Phe Gln Ala Ile Lys Thr His Ser Glu Ala Ser Gly Met Val Thr Gln
               405
Leu
<210> 69
```

```
<210> 69
<211> 2479
<212> DNA
<213> Homo sapien
<220>
<221> CDS
```

<222> (57)...(1535)

<223> Nucleotide sequence encoding human serine protease

<300>

<308> GenBank U75329 <309> 1997-10-10

<400> 69 gtcatattga acattccaga tacctatcat tactcgatgc tgttgataac agcaag atg 59 107 gct ttg aac tca ggg tca cca cca gct att gga cct tac tat gaa aac Ala Leu Asn Ser Gly Ser Pro Pro Ala Ile Gly Pro Tyr Tyr Glu Asn cat gga tac caa ccg gaa aac ccc tat ccc gca cag ccc act gtg gtc His Gly Tyr Gln Pro Glu Asn Pro Tyr Pro Ala Gln Pro Thr Val Val 155 ccc act gtc tac gag gtg cat ccg gct cag tac tac ccg tcc ccc gtg  $\mbox{Pro}$  Thr Val Tyr Glu Val  $\mbox{His}$   $\mbox{Pro}$  Ala Gln Tyr Tyr  $\mbox{Pro}$   $\mbox{Ser}$   $\mbox{Pro}$  Val 203 ccc cag tac gcc ccg agg gtc ctg acg cag gct tcc aac ccc gtc gtc Pro Gln Tyr Ala Pro Arg Val Leu Thr Gln Ala Ser Asn Pro Val Val 251 tgc acg cag ccc aaa tcc cca tcc ggg aca gtg tgc acc tca aag act 299 Cys Thr Gln Pro Lys Ser Pro Ser Gly Thr Val Cys Thr Ser Lys Thr 75 aag aaa gca ctg tgc atc acc ttg acc ctg ggg acc ttc ctc gtg gga 347 Lys Lys Ala Leu Cys Ile Thr Leu Thr Leu Gly Thr Phe Leu Val Gly gct gcg ctg gcc gct ggc cta ctc tgg aag ttc atg ggc agc aag tgc 395 Ala Ala Leu Ala Ala Gly Leu Leu Trp Lys Phe Met Gly Ser Lys Cys tcc aac tct ggg ata gag tgc gac tcc tca ggt acc tgc atc aac ccc 443 Ser Asn Ser Gly Ile Glu Cys Asp Ser Ser Gly Thr Cys Ile Asn Pro tct aac tgg tgt gat ggc gtg tca cac tgc ccc ggc ggg gag gac gag Ser Asn Trp Cys Asp Gly Val Ser His Cys Pro Gly Glu Asp Glu 491 aat cgg tgt gtt cgc ctc tac gga cca aac ttc atc ctt cag atg tac 539 Asn Arg Cys Val Arg Leu Tyr Gly Pro Asn Phe Ile Leu Gln Met Tyr 155 tca tct cag agg aag tcc tgg cac cct gtg tgc caa gac gac tgg aac 587 Ser Ser Gln Arg Lys Ser Trp His Pro Val Cys Gln Asp Asp Trp Asn 170 635 gag aac tac ggg cgg gcc tgc agg gac atg ggc tat aag aat aat Glu Asn Tyr Gly Arg Ala Ala Cys Arg Asp Met Gly Tyr Lys Asn Asn ttt tac tct agc caa gga ata gtg gat gac agc gga tcc acc agc ttt Phe Tyr Ser Ser Gln Gly Ile Val Asp Asp Ser Gly Ser Thr Ser Phe 683 195 205 atg aaa ctg aac aca agt gcc ggc aat gtc gat atc tat aaa aaa ctg 731 Met Lys Leu Asn Thr Ser Ala Gly Asn Val Asp Ile Tyr Lys Lys Leu

WO 01/57194		PCT/US01/03471
	61/66	

210					215					220					225	
	cac His															779
	gcc Ala															827
	ggt Gly															875
cac His	gtc Val 275	cag Gln	aac Asn	gtc Val	cac His	gtg Val 280	tgc Cys	gga Gly	ggc Gly	tcc Ser	atc Ile 285	atc Ile	acc Thr	ccc Pro	gag Glu	923
	atc Ile															971
	cat His															1019
	gga Gly															1067
	tcc Ser															1115
cct Pro	ctg Leu 355	act Thr	ttc Phe	aac Asn	gac Asp	cta Leu 360	gtg Val	aaa Lys	cca Pro	gtg Val	tgt Cys 365	ctg Leu	ccc Pro	aac Asn	cca Pro	1163
	atg Met															1211
gcc Ala	acc Thr	gag Glu	gag Glu	aaa Lys 390	Gly 999	aag Lys	acc Thr	tca Ser	gaa Glu 395	gtg Val	ctg Leu	aac Asn	gct Ala	gcc Ala 400	aag Lys	1259
	ctt Leu															1307
	ctg Leu															1355
	gat Asp 435															1403
	aat Asn															1451

•

Ala Lys Ala Tyr Arg Pro Gly Val Tyr Gly Asn Val Met Val Phe Thr 470  gac tgg att tat cga caa atg aag gca aac ggc taa tccacatggt  Asp Trp Ile Tyr Arg Gln Met Lys Ala Asn Gly 485  cttcgtcett gacgtcgttt tacaagaaaa caatggggct ggttttgctt ccccgtgcat gatttactct tagagatgat tcagaggtca cttcattttt attaaacagt gaacttgtct ggctttggca ctctctgcca tactgtgcag gctgcagtgg ctccctgcc cagcctgct tccctaaccc cttgtccga aggggtgatg gccggctggt tgtgggcact ggcggtcaat tgtggaagga agagggttgg aggctgccc cattgagatc ttcctgcta gctgggtcaat ttggatgaa gggagacagc cagtggcac ctgctggggggaagggggggggg																		
Asp Trp Ile Tyr Arg Gln Met Lys Ala Asn Gly * 485		gcc Ala	aaa Lys	gct Ala	tac Tyr	Arg	cca Pro	gga Gly	gtg Val	tac Tyr	Gly	aat Asn	gtg Val	atg Met	gta Val	Phe	acg Thr	1499
gatttactct tagagatgat tcagagtca cttcatttt attaacagt gaactttgct 1 ggctttggca ctctctgcca tactgtgag gctgcagtgg ctccctgcc cagcctgctc 1 tccctaaccc cttgtccca aggggtgatg gccggctggt tgtgggcact ggcggtcat 1 tgtggaagga aggaggttgg aggctgccc cattgagat ttctgctgag gcctttcca 1 ggggccaatt ttggatgagc atggagctgt cacttccag ctgctggatg acttgagatg 1 aaaaaggaga gcaggatgaa gggaagcagc caggtggcac ctgcagcggc tgccctctgg ggccacttg tagtgtccc agcctactc acaaggggat tttgctgatg ggttcttaga 2 gccttagcag ccctggatgg tggccagaaa taaagggacc agccttcat gggtggtgatc ctgtaagggg gacaagaaaac atttttgtt ttatggggg agaatataga 2 aggtgcactt ggtgcagagg aagcaagaaac atttttgtt ttatggggg agaatataga 2 aggtcccac ctgcaacttg ggtggggacc ctgggggggag accaagcact cctggtgc 2 aggtctcac ctgcaacttg ggtggggac ctggggggag agccattg cctgggaggag agccattg cgccaggttg gaccattg ggcctgtcc ggcctgtcc ggccagttg gccctggacg ctgcctctcac 2 cctccctgac ctgctccta gcacctgga gagtgaatgc ccttggtcc ctggagggga 2 cgcaagttg gcaccttg ggcctcttca ggcctgatag tcattggaa ttgaggcc 2 ggcaagtttg gcaccttga ttcaaagcac tctt  <210 > 70 <211 > 492 <212 > PRT <213 > Homo sapien  <400 > 70  Met Ala Leu Asn Ser Gly Ser Pro Pro Ala Ile Gly Pro Tyr Tyr Glu 1 1	2	gac Asp	tgg Trp	att Ile	Tyr	cga Arg	caa Gln	atg Met	aag Lys	Ala	aac Asn	ggc Gly	taa *	tcca	acat	ggt		1545
<pre>&lt;211&gt; 492 &lt;212&gt; PRT &lt;212&gt; Homo sapien </pre> <pre>&lt;400&gt; 70 Met Ala Leu Asn Ser Gly Ser Pro Pro Ala Ile Gly Pro Tyr Tyr Glu 1</pre>		gatt ggct tccc tgggg aaac gcct gcagt aggt accca gcca	ttact ttact ttaa ggaag gcaa gcact ttag gccc cctc cct	tot to go of the control of the cont	tagagettetetetetetetetetetetetetetetetet	gatga ctgcc ggtt ggtt gatga tcgag gtca ggat gacat tcct gatga gatga	at to a t	cagagactgactgggctgggggggggggggggggggggg	ggtcaggtcaggccccggccggacaggaaaaaaaaaaggaaaaaggaaaaggaaaaggaaaaggactccaggactccaggactcaggactcaagggctcaagggccccccc	a ctt go go cat cac acc acc acc acc acc acc acc acc	tcati tgcaç cggci ttgaç cttci ggtgg aaggg ttttç aggaa gggaç gtga cctga	ttt gtgg ggt gatc gac gac gttc gac gttc gac gttc gac gttc	atta ctco tgto ctgo ctgo ttto agco ttai gcco gaci ccci	aaaca ccctg gggca ctgga cagcg cctga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga ccaga cca cc	agt g gcc g act g atg a	gaact cagco ggcgg gtcct actt ggtt gggt ctcct ctct ct	ttgtct ctgctc gtcaat gttcaa gattgag cttaga gtgtga gatag ctcaggc caggca	1605 1665 1725 1785 1845 1905 2025 2085 2145 2205 2325 2385 2445 2479
Met Ala Leu Asn Ser Gly Ser Pro Pro Ala Ile Gly Pro Tyr Tyr Glu 1       15         Asn His Gly Tyr Gln Pro Glu Asn Pro Tyr Pro Ala Gln Pro Thr Val 20       25         Val Pro Thr Val Tyr Glu Val His Pro Ala Gln Tyr Tyr Pro Ser Pro 35       40         Val Pro Gln Tyr Ala Pro Arg Val Leu Thr Gln Ala Ser Asn Pro Val 50       80         Val Cys Thr Gln Pro Lys Ser Pro Ser Gly Thr Val Cys Thr Ser Lys 65       70         Thr Lys Lys Ala Leu Cys Ile Thr Leu Thr Leu Gly Thr Phe Leu Val 90       95         Gly Ala Ala Leu Ala Ala Gly Leu Leu Trp Lys Phe Met Gly Ser Lys 100       105         Cys Ser Asn Ser Gly Ile Glu Cys Asp Ser Ser Gly Thr Cys Ile Asn 115       125         Pro Ser Asn Trp Cys Asp Gly Val Ser His Cys Pro Gly Gly Glu Asp 130       135         Glu Asn Arg Cys Val Arg Leu Tyr Gly Pro Asn Phe Ile Leu Gln Met 145       150         Tyr Ser Ser Gln Arg Lys Ser Trp His Pro Val Cys Gln Asp Asp Trp 165       170         Asn Pro Gly Arg Ala Ala Cys Arg Asp Met Gly Tyr Lys Asn 180       185         Phe Met Lys Leu Asn Thr Ser Ala Cys Arg Asp Met Gly Tyr Lys Lys 210         Phe Met Lys Leu Asn Thr Ser Ala Cys Arg Asp Val Asp Ile Tyr Lys Lys 220         Phe Met Lys Leu Asn Thr Ser Ala Cys Ser Ser Lys Ala Val Val Ser Leu Arg 220		<211 <212	L> 49 2> PI	92 RT	sapie	en												
Asn His Gly Tyr Cln       Tyr Gln       Pro Glu       Asn Pro Tyr       Pro Ala Gln       Pro Thr Val 30         Val Pro Thr Val Tyr Gly       Val His Pro Ala Gln       Tyr Tyr Pro Ser Pro 30       Pro 40       Tyr Pro 45       Pro 45         Val Pro Gln       Tyr Ala Pro Arg Val Leu Thr Gln       Ala Ser Asn Pro Val 60       Val Cys Thr Ser Lys 60         Val Cys Thr Gln       Pro Lys Ser Pro Ser Gly Thr Val Cys Thr Ser Lys 70       80         Thr Lys Lys Lys Ala Leu Cys Ile Thr Leu Thr Leu Thr Leu Gly Thr Phe Leu Val 85       90         Gly Ala Ala Leu Ala Ala Gly Leu Leu Trp Lys Phe Met Gly Ser Lys 100       105         Cys Ser Asn Ser Gly Ile Glu Cys Asp Ser Ser Gly Thr Cys Ile Asn 115       120         Pro Ser Asn Trp Cys Asp Gly Val Ser His Cys Pro Asn Phe Ile Leu Gln Met 145       150         Glu Asn Arg Cys Val Arg Leu Tyr Gly Pro Asn Phe Ile Leu Gln Met 145       150         Tyr Ser Ser Gln Arg Lys Ser Trp His Pro 165       170         Asn Phe Tyr Gly Arg Ala Ala Cys Asp Arg Asp Met Gly Tyr Lys Asn 180         Asn Phe Tyr Gly Arg Ala Ala Cys Arg Arg Asp Ser Gly Ser Thr Ser 195         Phe Met Lys Leu Asn Thr Ser Ala Gly Asn Val Asp Ile Tyr Lys Lys 210         Leu Tyr His Ser Asp Ala Cys Ser Ser Lys Ala Val Val Ser Leu Arg 225		Met			Asn	_	Gly	Ser	Pro	Pro		Ile	Gly	Pro	Tyr	_	Glu	
Val       Pro       Thr       Val       Tyr       Glu       Val       His       Pro       Ala       Gln       Tyr       Pro       Ser       Pro       45         Val       Pro       Gln       Tyr       Ala       Pro       Arg       Val       Leu       Thr       Gln       Ala       Ser       Pro       Val       Gln       Ala       Ser       Lys       Ala       Leu       Lys       Ser       Gly       Thr       Val       Cys       Thr       Ser       Lys       Ala       Leu       Cys       Ile       Thr       Leu       Thr       Val       Cys       Thr       Pro       Ser       Lys       Ala       Leu       Cys       Ile       Thr       Leu       Thr       Val       Cys       Thr       Pro       Leu       Val       Ser       Lys       Pro       Leu       Cys       Thr       Pro       He       Met       Gly       Ser       Lys	2		His	Gly		_	Pro	Glu	Asn			Pro	Ala	Gln			Val	
Val       Pro       Gln       Tyr       Ala       Pro       Arg       Val       Leu       Thr       Gln       Ala       Ser       Pro       Val       Cys       Thr       Ser       Lys       Ser       Pro       Ser       Pro       Ser       Gly       Thr       Leu       Cys       Thr       Val       Cys       Thr       Ser       Lys       Bo         Thr       Lys       Lys       Ala       Leu       Cys       Ile       Thr       Leu       Thr       Leu       Gly       Thr       Phe       Leu       Val         Gly       Ala       Ala       Ala       Gly       Leu       Leu       Trp       Lys       Phe       Met       Gly       Ser       Lys         Pro       Ser       Asn       Ser       Gly       Ile       Glu       Cys       Asp       Ser       Ser       Gly       Ile       Asn       Ile       Ile       Asn       Ile       Il	٦	Val	Pro			Tyr	Glu	Val			Ala	Gln	Tyr			Ser	Pro	
Val Cys         Thr Gln         Pro         Lys         Ser         Pro         Ser         Gly         Thr Val         Cys         Thr Ser         Lys         80           Thr Lys         Lys         Ala         Leu         Cys         Ile         Thr         Leu         Thr         Leu         Gly         Thr         Phe         Leu         Val         Phe         Met         Gly         Ser         Lys         Lys         Lys         Phe         Met         Gly         Ser         Lys         Lys         Lys         Phe         Met         Gly         Ser         Lys	,	Val	_		Tyr	Ala	Pro	_		Leu	Thr	Gln			Asn	Pro	Val	
Thr Lys Lys Ala Leu Cys Ile Thr Leu Thr Leu Gly Thr Phe Leu Val 85 Gly Ala Ala Leu Ala Ala Gly Leu Leu Trp Lys Phe Met Gly Ser Lys 100 Cys Ser Asn Ser Gly Ile Glu Cys Asp Ser Ser Gly Thr Cys Ile Asn 115 Pro Ser Asn Trp Cys Asp Gly Val Ser His Cys Pro Gly Gly Glu Asp 130 Glu Asn Arg Cys Val Arg Leu Tyr Gly Pro Asn Phe Ile Leu Gln Met 145 Tyr Ser Ser Gln Arg Lys Ser Trp His Pro Val Cys Gln Asp Asp Trp 165 Asn Glu Asn Tyr Gly Arg Ala Ala Cys Arg Asp Asp Asp Met Gly Tyr Lys Asn 180 Asn Phe Tyr Ser Ser Gln Gly Ile Val Asp Asp Asp Ser Gly Ser Thr Ser 195 Phe Met Lys Leu Asn Thr Ser Ala Gly Asn Val Asp Ile Tyr Lys Lys 210 Leu Tyr His Ser Asp Ala Cys Ser Ser Lys Ala Val Val Ser Leu Arg 225 Leu Tyr His Ser Asp Ala Cys Ser Ser Lys Ala Val Val Ser Leu Arg 225				Thr	Gln	Pro	-		Pro	Ser	Gly			Cys	Thr	Ser	-	
Gly Ala Ala Leu Ala Ala Gly Leu Leu Trp Lys Phe Met Gly Ser Lys			Lys	Lys	Ala			Ile	Thr	Leu			Gly	Thr	Phe			
Cys       Ser       Asn       Ser       Gly       Ile       Glu       Cys       Asp       Ser       Ser       Gly       Thr       Cys       Ile       Asn         Pro       Ser       Asn       Trp       Cys       Asp       Gly       Val       Ser       His       Cys       Pro       Gly       Gly       Gly       Asp         Glu       Asn       Arg       Cys       Val       Arg       Leu       Tyr       Gly       Pro       Asn       Phe       Ile       Leu       Gln       Met         Tyr       Ser       Ser       Gln       Leu       Tyr       Gly       Pro       Asn       Phe       Ile       Leu       Gln       Met         Tyr       Ser       Ser       Gln       Leu       Tyr       His       Pro       Val       Cys       Gln       Asp       Asp       Tyr       Lys       Asp       Trp       Into       Into <t< td=""><td>(</td><td>Gly</td><td>Ala</td><td>Ala</td><td></td><td></td><td>Ala</td><td>Gly</td><td>Leu</td><td></td><td></td><td>Lys</td><td>Phe</td><td>Met</td><td></td><td></td><td>Lys</td><td></td></t<>	(	Gly	Ala	Ala			Ala	Gly	Leu			Lys	Phe	Met			Lys	
Pro       Ser       Asn       Trp       Cys       Asp       Gly       Val       Ser       His       Cys       Pro       Gly       Gly       Gly       Asp       140       140       140       140       140       140       140       140       140       140       140       140       140       140       140       140       140       140       140       140       140       140       140       140       140       160       160       160       160       160       160       160       160       160       160       160       160       160       160       160       160       160       170       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       175       1	(	Cys	Ser			Gly	Ile	Glu			Ser	Ser	Gly			Ile	Asn	
Glu Asn Arg Cys Val Arg Leu Tyr Gly Pro Asn Phe Ile Leu Gln Met 145		Pro			Trp	Cys	Asp			Ser	His	Cys			Gly	Glu	Asp	
Tyr Ser Ser Gln Arg Lys Ser Trp His Pro Val Cys Gln Asp Asp Trp 165				Arg	Cys	Val			Tyr	Gly	Pro			Ile	Leu	Gln		
Asn Glu Asn Tyr Gly Arg Ala Ala Cys Arg Asp Met Gly Tyr Lys Asn 180			Ser	Ser	Gln			Ser	Trp	His			Cys	Gln	Asp	-		
Asn Phe Tyr Ser Ser Gln Gly Ile Val Asp Asp Ser Gly Ser Thr Ser  195	2	Asn	Glu	Asn			Arg	Ala	Ala	Cys		Asp	Met	Gly	Tyr		Asn	
Phe Met Lys Leu Asn Thr Ser Ala Gly Asn Val Asp Ile Tyr Lys Lys 210 215 220  Leu Tyr His Ser Asp Ala Cys Ser Ser Lys Ala Val Val Ser Leu Arg 225 230 235 240	ž	Asn	Phe			Ser	Gln	Gly			Asp	Asp	Ser			Thr	Ser	
Leu Tyr His Ser Asp Ala Cys Ser Ser Lys Ala Val Ser Leu Arg 225 230 235 240	1	Phe			Leu	Asn	Thr			Gly	Asn	Val			Tyr	Lys	Lys	
				His	Ser	Asp			Ser	Ser	Lys			Val	Ser	Leu		
Cyp Deu Aid Cyp Giy vai Ash Deu Ash Sel Sel Ald Gill Sel Ard lie			T.e.i	7/1-	C17C	٠, ٢		7\ ~~~	T 03-	7\ ~~	C~~		7\ ~~~	~TD	C~~	7\ ~~~		
	(	Cys	пеп	нта	CAR	GTĀ	vaT	ASII	ьеu	ASII	ser	ser	Arg	GTII	ser	ALG	TTG	

```
245
                                      250
Val Gly Gly Glu Ser Ala Leu Pro Gly Ala Trp Pro Trp Gln Val Ser
            260
                                 265
                                                      270
Leu His Val Gln Asn Val His Val Cys Gly Gly Ser Ile Ile Thr Pro
                             280
        275
                                                  285
Glu Trp Ile Val Thr Ala Ala His Cys Val Glu Lys Pro Leu Asn Asn
    290
                         295
                                              300
Pro Trp His Trp Thr Ala Phe Ala Gly Ile Leu Arg Gln Ser Phe Met
                     310
                                          315
Phe Tyr Gly Ala Gly Tyr Gln Val Gln Lys Val Ile Ser His Pro Asn
                 325
                                      330
Tyr Asp Ser Lys Thr Lys Asn Asn Asp Ile Ala Leu Met Lys Leu Gln
            340
                                 345
                                                       350
Lys Pro Leu Thr Phe Asn Asp Leu Val Lys Pro Val Cys Leu Pro Asn
        355
                             360
                                                  365
Pro Gly Met Met Leu Gln Pro Glu Gln Leu Cys Trp Ile Ser Gly Trp
    370
                         375
                                              380
Gly Ala Thr Glu Glu Lys Gly Lys Thr Ser Glu Val Leu Asn Ala Ala
                     390
                                          395
Lys Val Leu Leu Ile Glu Thr Gln Arg Cys Asn Ser Arg Tyr Val Tyr
                 405
                                      410
                                                           415
Asp Asn Leu Ile Thr Pro Ala Met Ile Cys Ala Gly Phe Leu Gln Gly
                                  425
            420
Asn Val Asp Ser Cys Gln Gly Asp Ser Gly Gly Pro Leu Val Thr Ser
                             440
Asn Asn Asn Ile Trp Trp Leu Ile Gly Asp Thr Ser Trp Gly Ser Gly
                         455
                                              460
Cys Ala Lys Ala Tyr Arg Pro Gly Val Tyr Gly Asn Val Met Val Phe
                     470
                                         475
Thr Asp Trp Ile Tyr Arg Gln Met Lys Ala Asn Gly
                 485
<210> 71
<211> 2079
<212> DNA
<213> Homo sapien
<220>
<221> CDS
<222> (251)...(1522)
<223> Nucleotide sequence encoding transmembrane
      protease, serine 4 (TMPRSS4)
<300>
<308> GenBank NM016425
<309> 2000-11-06
<400> 71
gagaggcagc agettgttca geggacaagg atgetgggeg tgagggacca aggeetgeec
tgeacteggg cetectecag ccagtgetga ccagggaett etgacetget ggecagecag
                                                                        120
gacctgtgtg gggaggccct cctgctgcct tggggtgaca atctcagctc caggctacag
                                                                         180
ggagaccggg aggatcacag agccagcatg gtacaggatc ctgacagtga tcaacctctg
                                                                        240
aacageeteg atg tea aac eee tge gea aac eee gta tee eea tgg aga
                                                                        289
            Met Ser Asn Pro Cys Ala Asn Pro Val Ser Pro Trp Arg
cct tca gaa agt gtg ggg atc ccc atc atc ata gca cta ctg agc ctg
                                                                         337
Pro Ser Glu Ser Val Gly Ile Pro Ile Ile Ile Ala Leu Leu Ser Leu
     15
gcg agt atc atc att gtg gtt gtc ctc atc aag gtg att ctg gat aaa Ala Ser Ile Ile Ile Val Val Leu Ile Lys Val Ile Leu Asp Lys
                                                                        385
```

30					35					40					45	
					Gly 999											433
					ctg Leu											481
					ccc Pro											529
					ctg Leu											577
					gac Asp 115											625
					tac Tyr											673
					gat Asp											721
cag Gln	gag Glu	ctt Leu 160	cgc Arg	atg Met	cgg Arg	aac Asn	tca Ser 165	agt Ser	Gly 333	ccc Pro	tgt Cys	ctc Leu 170	tca Ser	ggc Gly	tcc Ser	769
					tgt Cys											817
					gag Glu 195											865
					gac Asp											913
gac Asp	ccc Pro	cac His	tgg Trp 225	gtc Val	ctc Leu	acg Thr	gca Ala	gcc Ala 230	cac His	tgc Cys	ttc Phe	agg Arg	aaa Lys 235	cat His	acc Thr	961
					aag Lys											1009
ttc Phe	cca Pro 255	tcc Ser	ctg Leu	gct Ala	gtg Val	gcc Ala 260	aag Lys	atc Ile	atc Ile	atc Ile	att Ile 265	gaa Glu	ttc Phe	aac Asn	ccc Pro	1057
					aat Asn 275											1105

ctc act ttc tca ggc aca gtc agg ccc atc tgt ctg ccc ttc ttt gat Leu Thr Phe Ser Gly Thr Val Arg Pro Ile Cys Leu Pro Phe Phe Asp 290 295 300	1153								
gag gag ctc act cca gcc acc cca ctc tgg atc att gga tgg ggc ttt Glu Glu Leu Thr Pro Ala Thr Pro Leu Trp Ile Ile Gly Trp Gly Phe 305 310 315	1201								
acg aag cag aat gga ggg aag atg tct gac ata ctg ctg cag gcg tca Thr Lys Gln Asn Gly Gly Lys Met Ser Asp Ile Leu Leu Gln Ala Ser 320 325 330	1249								
gtc cag gtc att gac agc aca cgg tgc aat gca gac gat gcg tac cag Val Gln Val Ile Asp Ser Thr Arg Cys Asn Ala Asp Asp Ala Tyr Gln 335 340 345	1297								
ggg gaa gtc acc gag aag atg atg tgt gca ggc atc ccg gaa ggg ggt Gly Glu Val Thr Glu Lys Met Met Cys Ala Gly Ile Pro Glu Gly Gly 350 355 360 365	1345								
gtg gac acc tgc cag ggt gac agt ggt ggg ccc ctg atg tac caa tct Val Asp Thr Cys Gln Gly Asp Ser Gly Gly Pro Leu Met Tyr Gln Ser 370 375 380	1393								
gac cag tgg cat gtg gtg ggc atc gtt agc tgg ggc tat ggc tgc ggg Asp Gln Trp His Val Val Gly Ile Val Ser Trp Gly Tyr Gly Cys Gly 385 390 395	1441								
ggc ccg agc acc cca gga gta tac acc aag gtc tca gcc tat ctc aac Gly Pro Ser Thr Pro Gly Val Tyr Thr Lys Val Ser Ala Tyr Leu Asn 400 405 410	1489								
tgg atc tac aat gtc tgg aag gct gag ctg taa tgctgctgcc cctttgcagt Trp Ile Tyr Asn Val Trp Lys Ala Glu Leu * 415 420	1542								
gctgggagcc gcttccttcc tgccctgccc acctggggat ccccaaagt cagacacaga gcaagagtcc ccttgggtac accctctgc ccacagctc agcattctt ggagcagcaa agggcctcaa ttcctgtaag agaccctcgc agcccagagg cgcccagagg aagtcagcag ccctagctcg gccacacttg gtgctcccag catcccaggg agagacacag ccactgaac aaggtctcag gggtattgct aagccaagaa ggaccttcc cacactactg aatggaagca ggctgtcttg taaaagccca gatcactgtg ggctggagag gagaaggaaa gggtctgcgc cagccctgtc cgtcttcacc catccccaag cctactagag caagaaacca gttgtaatat aaaatgcact gccctactgt tggtatgact accgttacct actgttgtca ttgttattac agctatggcc actattatta aagagctgtg taacatcaaa aaaaaaaaaa									
<210> 72 <211> 423 <212> PRT <213> Homo sapien									
<pre>&lt;400&gt; 72 Met Ser Asn Pro Cys Ala Asn Pro Val Ser Pro Trp Arg Pro Ser Glu 1 5 10 15</pre>									
Ser Val Gly Ile Pro Ile Ile Ile Ala Leu Leu Ser Leu Ala Ser Ile									
20 25 30  Ile Ile Val Val Leu Ile Lys Val Ile Leu Asp Lys Tyr Tyr Phe 35 40 45									
Leu Cys Gly Gln Pro Leu His Phe Ile Pro Arg Lys Gln Leu Cys Asp 50 55 60									
Gly Glu Leu Asp Cys Pro Leu Gly Glu Asp Glu Glu His Cys Val Lys 65 70 75 80									
•									

Ser	Phe	Pro	Glu	Gly 85	Pro	Ala	Val	Ala	Val 90	Arg	Leu	Ser	Lys	Asp 95	Arg
Ser	Thr	Leu	Gln 100	Val	Leu	Asp	Ser	Ala 105	Thr	Gly	Asn	Trp	Phe 110	Ser	Ala
Cys	Phe	Asp 115	Asn	Phe	Thr	Glu	Ala 120	Leu	Ala	Glu	Thr	Ala 125	Cys	Arg	Gln
Met	Gly 130	Tyr	Ser	Ser	Lys	Pro 135	Thr	Phe	Arg	Ala	Val 140	Glu	Ile	Gly	Pro
145		-		_	150					155				Glu	160
				165					170					Val 175	
			180			_	_	185		_			190	Val	
-	-	195					200		_		_	205		Ser	
	210					215	-	_			220			Pro	
225					230		_		_	235			_	Val	240
				245		_		_	250					Pro 255	
			260	_				265					270	Tyr	
_	_	275	_				280	-				285		Thr	
	290			_		295	_				300	-		Glu	
305					310	_				315				Lys	320
	_	_	-	325		_			330					Gln 335	
	_		340	_	_			345	_		_		350	Glu	
		355			-	•	360				_	365		Asp	
	370					375					380			Gln	
385					390					395				Pro	400
				405		_	Val	Ser	Ala 410	Tyr	Leu	Asn	Trp	Ile 415	Tyr
Asn	Val	Trp	Lys 420	Ala	Glu	Leu									